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(57) Abstract

This invention relates to plant fatty acid hydroxylases. Methods to use conserved amino acid or nucleotide sequences to obtain plant fatty acid hydroxylases are described. Also described is the use of cDNA clones encoding a plant hydroxylase to produce a family of hydroxylated fatty acids in transgenic plants. In addition, the use of genes encoding fatty acid hydroxylases or desaturases to alter the level of lipid fatty acid unsaturation in transgenic plants is described.

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PRODUCTION OF HYDROXYLATED FATTY ACIDS IN GENETICALLY MODIFIED PLANTS

TECHNICAL FIELD

The present invention concerns the

identification of nucleic acid sequences and
constructs, and methods related thereto, and the use
of these sequences and constructs to produce
genetically modified plants for the purpose of
altering the fatty acid composition of plant oils,
waxes and related compounds.

DEFINITIONS

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The subject of this invention is a class of enzymes that introduce a hydroxyl group into several different fatty acids resulting in the production of several different kinds of hydroxylated fatty acids. 15 In particular, these enzymes catalyze hydroxylation of oleic acid to 12-hydroxy oleic acid and icosenoic acid to 14-hydroxy icosenoic acid. Other fatty acids such as palmitoleic and erucic acids may also be substrates. Since it is not possible to refer to the 20 enzyme by reference to a unique substrate or product, the enzyme is referred throughout as kappa hydroxylase to indicate that the enzyme introduces the hydroxyl three carbons distal (i.e., away from 25 the carboxyl carbon of the acyl chain) from a double bond located near the center of the acyl chain.

The following fatty acids are also the subject of this invention: ricinoleic acid, 12-hydroxyoctadec-cis-9-enoic acid (120H-18:1cisas); lesquerolic acid, 14-hydroxy-cis-11-icosenoic acid (140H-20:1cisaii); densipolic acid, 12-hydroxyoctadec-cis-9,15-dienoic acid (120H-18:2cisas,15); auricolic acid, 14-hydroxy-cis-11,17-icosadienoic acid (140H-

20:2cisall.17); hydroxyerucic, 16-hydroxydocos-cis-13-enoic acid (160H-22:1cisall); hydroxypalmitoleic, 12-hydroxyhexadec-cis-9-enoic (120H-16:1cisall); icosenoic acid (20:1cisall). It will be noted that icosenoic acid is spelled eicosenoic acid in some countries.

BACKGROUND

Extensive surveys of the fatty acid composition of seed oils from different species of higher plants have resulted in the identification of at least 33 structurally distinct monohydroxylated 10 plant fatty acids, and 12 different polyhydroxylated fatty acids that are accumulated by one or more plant species (reviewed by van de Loo et al., 1993). Ricinoleic acid, the principal constituent of the seed oil from the castor plant Ricinus communis 15 (L.), is of commercial importance. The present inventors have cloned a gene from this species that encodes a fatty acid hydroxylase, and have used this gene to produce ricinoleic acid in transgenic plants 20 of other species. Some of this scientific evidence has been published by the present inventors (van de Loo et al., 1995).

The use of the castor hydroxylase gene to also produce other hydroxylated fatty acids such as lesquerolic acid, densipolic acid, hydroxypalmitoleic, hydroxyerucic and auricolic acid in transgenic plants is the subject of this invention. In addition, the identification of a gene encoding a homologous hydroxylase from Lesquerella fendleri, and the use of this gene to produce these hydroxylated fatty acids in transgenic plants is the subject of this invention.

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Castor is a minor oilseed crop. Approximately 50% of the seed weight is oil (triacylglycerol) in which 85-90% of total fatty acids are the hydroxylated fatty acid, ricinoleic acid. Oil pressed or extracted from castor seeds has many industrial uses based upon the properties endowed by the hydroxylated fatty acid. The most important uses are production of paints and varnishes, nylon-type synthetic polymers, resins, lubricants, and cosmetics (Atsmon, 1989).

In addition to oil, the castor seed contains the extremely toxic protein ricin, allergenic proteins, and the alkaloid ricinine. These constituents preclude the use of the untreated seed meal (following oil extraction) as a livestock feed, normally an important economic aspect of oilseed utilization. Furthermore, with the variable nature of castor plants and a lack of investment in breeding, castor has few favorable agronomic characteristics.

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For a combination of these reasons, castor is no longer grown in the United States and the development of an alternative domestic source of hydroxylated fatty acids would be attractive. The production of ricinoleic acid, the important constituent of castor oil, in an established oilseed crop through genetic engineering would be a particularly effective means of creating a domestic source.

Because there is no practical source of lesquerolic, densipolic and auricolic acids from plants that are adapted to modern agricultural practices, there is currently no large-scale use of these fatty acids by industry. However, the fatty

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acids would have uses similar to those of ricinoleic acid if they could be produced in large quantities at comparable cost to other plant-derived fatty acids (Smith, 1985).

Plant species, such as certain species in the genus Lesquerella, that accumulate a high proportion of these fatty acids, have not been domesticated and are not currently considered a practical source of fatty acids (Hirsinger, 1989). This invention represents a useful step toward the eventual production of these and other hydroxylated fatty acids in transgenic plants of agricultural importance.

The taxonomic relationships between plants having similar or identical kinds of unusual fatty 15 acids have been examined (van de Loo et al., 1993). In some cases, particular fatty acids occur mostly or solely in related taxa. In other cases there does not appear to be a direct link between taxonomic relationships and the occurrence of unusual fatty 20 acids. In this respect, ricinoleic acid has now been identified in 12 genera from 10 families (reviewed in van de Loo et al., 1993). Thus, it appears that the ability to synthesize hydroxylated fatty acids has evolved several times independently during the 25 radiation of the angiosperms. This suggested to us that the enzymes which introduce hydroxyl groups into fatty acids arose by minor modifications of a related enzyme.

Indeed, as shown herein, the sequence similarity between A12 fatty acid desaturases and the kappa hydroxylase from castor is so high that it is not possible to unambiguously determine whether a particular enzyme is a desaturase or a hydroxylase

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on the basis of evidence in the scientific literature. Similarly, a patent application (PCT WO 94/11516) that purports to teach the isolation and use of Δ 12 fatty acid desaturases does not teach how to distinguish a hydroxylase from a desaturase. In 5 view of the importance of being able to distinguish between these activities for the purpose of genetic engineering of plant oils, the utility of that application is limited to the several instances where direct experimental evidence (e.g., altered fatty acid composition in transgenic plants) was presented to support the assignment of function. A method for distinguishing between fatty acid desaturases and fatty acid hydroxylases on the basis of amino acid sequence of the enzyme is also a subject of this invention.

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A feature of hydroxylated or other unusual fatty acids is that they are generally confined to seed triacylglycerols, being largely excluded from 20 the polar lipids by unknown mechanisms (Battey and Ohlrogge 1989; Prasad et al., 1987). This is particularly intriguing since diacylglycerol is a precursor of both triacylglycerol and polar lipid. With castor microsomes, there is some evidence that the pool of ricinoleoyl-containing polar lipid is 25 minimized by a preference of diacylglycerol acyltransferase for ricinoleate-containing diacylglycerols (Bafor et al., 1991). Analyses of vegetative tissues have generated few reports of 30 unusual fatty acids, other than those occurring in the cuticle. The cuticle contains various hydroxylated fatty acids which are interesterified to produce a high molecular weight polyester which serves a structural role. A small number of other

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exceptions exist in which unusual fatty acids are found in tissues other than the seed.

The biosynthesis of ricinoleic acid from oleic acid in the developing endosperm of castor (Ricinus communis) has been studied by a variety of methods. Morris (1967) established in double-labeling studies that hydroxylation occurs directly by hydroxyl substitution rather than via an unsaturated-, keto- or epoxy-intermediate.

Hydroxylation using oleoyl-CoA as precursor can be demonstrated in crude preparations or microsomes, but activity in microsomes is unstable and variable, and isolation of the microsomes involved a considerable, or sometimes complete loss of activity

(Galliard and Stumpf, 1966; Moreau and Stumpf, 1981). Oleic acid can replace oleoyl-CoA as a precursor, but only in the presence of CoA, Mg² and ATP (Galliard and Stumpf, 1966) indicating that activation to the acyl-CoA is necessary. However, no radioactivity could be detected in ricinoleoyl-CoA

radioactivity could be detected in ricinoleoyl-CoA (Moreau and Stumpf, 1981). These and more recent observations (Bafor et al., 1991) have been interpreted as evidence that the substrate for the castor oleate hydroxylase is oleic acid esterified to phosphatidylcholine or another phospholipid.

The hydroxylase is sensitive to cyanide and azide, and dialysis against metal chelators reduces activity, which could be restored by addition of FeSO₄, suggesting iron involvement in enzyme activity (Galliard and Stumpf, 1966). Ricinoleic acid synthesis requires molecular oxygen (Galliard and Stumpf, 1966; Moreau and Stumpf 1981) and requires NAD(P)H to reduce cytochrome b5 which is thought to be the intermediate electron donor for the

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hydroxylase reaction (Smith et al., 1992). Carbon monoxide does not inhibit hydroxylation, indicating that a cytochrome P450 is not involved (Galliard and Stumpf, 1966; Moreau and Stumpf 1981). Data from a study of the substrate specificity of the hydroxylase show that all substrate parameters (i.e., chain length and double bond position with respect to both ends) are important; deviations in these parameters caused reduced activity relative to oleic acid (Howling et al., 1972). The position at which the hydroxyl was introduced, however, was determined by the position of the double bond, always being three carbons distal. Thus, the castor acyl hydroxylase enzyme can produce a family of different hydroxylated fatty acids depending on the availability of substrates. Thus, as a matter of convenience, the enzyme is referred throughout this specification as a kappa hydroxylase (rather than an oleate hydroxylase) to indicate the broad substrate specificity.

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The castor kappa hydroxylase has many superficial similarities to the microsomal fatty acyl desaturases (Browse and Somerville, 1991). In particular, plants have a microsomal oleate desaturase active at the Δ12 position. The substrate of this enzyme (Schmidt et al., 1993) and of the hydroxylase (Bafor et al., 1991) appears to be a fatty acid esterified to the sn-2 position of phosphatidylcholine. When oleate is the substrate, the modification occurs at the same position (Δ12) in the carbon chain, and requires the same cofactors, namely electrons from NADH via cytochrome b₅ and molecular oxygen. Neither enzyme is inhibited

by carbon monoxide (Moreau and Stumpf, 1981), the characteristic inhibitor of cytochrome P450 enzymes.

There do not appear to have been any published biochemical studies of the properties of the hydroxylase enzyme(s) in Lesquerella.

Conceptual basis of the invention

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The present inventors have described the use of a cDNA clone from castor for the production of ricinoleic acid in transgenic plants. As noted above, biochemical studies had suggested that the 10 castor hydroxylase may not have strict specificity for oleic acid but would also catalyze hydroxylation of other fatty acids such as icosenoic acid $(20:1^{cisall})$ (Howling et al., 1972). Based on these studies, expression of kappa hydroxylase in 15 transgenic plants of species such as Brassica napus and Arabidopsis thaliana that accumulate fatty acids such as icosenoic acid (20:1cisall) and erucic acid (13-docosenoic acid; 22:1cisal3) may cause the 20 accumulation of hydroxylated derivatives of these fatty acids due to the activity of the hydroxylase on these fatty acids. Direct evidence is presented in Example 1 that hydroxlyated derivatives of ricinoleic, lesquerolic, densipolic and auricolic 25 fatty acids are produced in transgenic Arabidopsis plants.

Example 2 shows the isolation of a novel kappa hydroxylase gene from Lesquerella fendleri.

In view of the high degree of sequence similarity between $\Delta 12$ fatty acid desaturases and the castor hydroxylase (van de Loo et al., 1995), the validity of claims (e.g., PCT WO 94/11516) for using a limited set of desaturase or hydroxylase

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al., 1993).

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genes or sequences derived therefrom to identify genes of identical function from other species must be viewed with skepticism. In this application, the present inventors teach a method by which hydroxylase genes can be distinguished from desaturases. The present inventors describe a mechanistic basis for the similar reaction mechanisms of desaturases and hydroxylases. Briefly, the available evidence suggests that fatty acid desaturases have a similar reaction mechanism to the bacterial enzyme methane monooxygenase which catalyses a reaction involving oxygen-atom transfer $(CH_4 \rightarrow CH_3OH)$ (van de Loo et al., 1993). The cofactor in the hydroxylase component of methane monooxygenase is termed a μ -oxo bridged diiron cluster (FeOFe). The two iron atoms of the FeOFe cluster are liganded by protein-derived nitrogen or oxygen atoms, and are tightly redox-coupled by the covalently-bridging oxygen atom. The FeOFe cluster accepts two electrons, reducing it to the diferrous state, before oxygen binding. Upon oxygen binding, it is likely that heterolytic cleavage also occurs, leading to a high valent oxoiron reactive species that is stabilized by resonance rearrangements possible within the tightly coupled FeOFe cluster. The stabilized high-valent oxoiron state of methane monooxygenase is capable of proton extraction from methane, followed by oxygen transfer, giving methanol. The FeOFe cofactor has been shown to be directly relevant to plant fatty acid modifications by the demonstration that castor stearoyl-ACP

desaturase contains this type of cofactor (Fox et

On the basis of the foregoing considerations, the present inventors suggest that the castor oleate hydroxylase might be a structurally modified fatty acyl desaturase, based upon three arguments. The first argument involves the taxonomic distribution 5 of plants containing ricinoleic acid. Ricinoleic acid has been found in 12 genera of 10 families of higher plants (reviewed in van de Loo et al., 1993). Thus, plants in which ricinoleic acid occurs are found throughout the plant kingdom, yet close 10 relatives of these plants do not contain the unusual fatty acid. This pattern suggests that the ability to synthesize ricinoleic acid has arisen (and been lost) several times independently, and is therefore has recently diverged. In other words, the ability 15 to synthesize ricinoleic acid has evolved rapidly, suggesting that a relatively minor genetic change in the structure of the ancestral enzyme was necessary to accomplish it.

The second argument is that many biochemical properties of castor kappa hydroxylase are similar to those of the microsomal desaturases, as discussed above (e.g., both preferentially act on fatty acids esterified to the sn-2 position of

phosphatidylcholine, both use cytochrome b5 as an intermediate electron donor, both are inhibited by cyanide, both require molecular oxygen as a substrate, both are thought to be located in the endoplasmic reticulum).

The third argument stems from the discussion of oxygenase cofactors above, in which it is suggested that the plant membrane bound fatty acid desaturases may have a μ -oxo bridged diiron cluster-type cofactor, and that such cofactors are capable

of catalyzing both fatty acid desaturations and hydroxylations, depending upon the electronic and structural properties of the protein active site.

Taking these three arguments together, the present inventors suggest that kappa hydroxylase of 5 castor endosperm is homologous to the microsomal oleate $\Delta 12$ desaturase found in all plants. A number of genes encoding microsomal $\Delta 12$ desaturases from various species have recently been cloned (Okuley et al., 1994) and substantial information about the 10 structure of these enzymes is now known (Shanklin et al., 1994). Hence, in the following invention, the present inventors teach how to use structural information to isolate and identify kappa 15 hydroxylase genes. This example teaches the method by which any carbon-monoxide insensitive plant fatty acyl hydroxylase gene can be identified by one skilled in the art.

An unpredicted outcome of our studies on the 20 castor hydroxylase gene in transgenic Arabidopsis plants was the discovery that expression of the hydroxylase leads to increased accumulation of oleic acid in seed lipids. Because of the low nucleotide sequence homology between the castor hydroxylase and the $\Delta 12$ -desaturase (about 67%), it is unlikely that 25 this effect is due to silencing (also called sensesuppression or cosuppression) of the expression of the desaturase gene by the hydroxylase gene. Whatever the basis for the effect, this invention teaches the use of hydroxylase genes to alter the 30 level of fatty acid unsaturation in transgenic plants. This invention also teaches the use of genetically modified hydroxylase and desaturase

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genes to achieve directed modification of fatty acid unsaturation levels.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-D show the mass spectra of hydroxy fatty acids standards (Figure 1A, O-TMSmethylricinoleate; Figure 1B, O-TMS-methyl densipoleate; Figure 1C, O-TMS-methyl-lesqueroleate; and Figure 1D, O-TMS-methylauricoleate).

Figure 2 shows the fragmentation pattern of trimethylsilylated methyl esters of hydroxy fatty 10 acids.

Figure 3A shows the gas chromatogram of fatty acids extracted from seeds of wild type Arabidopsis plants. Figure 3B shows the gas chromatogram of fatty acids extracted from seeds of transgenic Arabidopsis plants containing the fahl2 hydroxylase gene. The numbers indicate the following fatty acids: [1] 16:0; [2] 18:0; [3] 18:1cisΔ9; [4] $18:2^{\text{cisa9.12}};$ [5] 20:0; [6] $20:1^{\text{cisa11}};$ [7] $18:3^{\text{cisa9.12.15}};$ [8] 20:2cisall.14; [9] 22:1clsall; [10] ricinoleic acid; [11] densipolic acid; [12] lesquerolic acid; and

[13] auricolic acid. Figures 4A-D show the mass spectra of novel

fatty acids found in seeds of transgenic plants. 25 Figure 4A shows the mass spectrum of peak 10 from Figure 3B. Figure 4B shows the mass spectrum of peak 11 from Figure 3B. Figure 4C shows the mass spectrum of peak 12 from Figure 3B. Figure 4D shows the mass spectrum of peak 13 from Figure 3B.

30 Figure 5 shows the nucleotide sequence of pLesg2 (SEQ ID NO:1).

Figure 6 shows the nucleotide sequence of pLesq3 (SEQ ID NO:2).

Figure 7 shows a Northern blot of total RNA from seeds of *L. fendleri* probed with pLesq2 or pLesq3. S, indicates RNA is from seeds; L, indicates RNA is from leaves.

Figures 8A-B show the nucleotide sequence of genomic clone encoding pLesq-HYD (SEQ ID NO:3), and the deduced amino acid sequence of hydroxylase enzyme encoded by the gene (SEQ ID NO:4).

Figures 9A-B show multiple sequence alignment
of deduced amino acid sequences for kappa
hydroxylases and microsomal Δ12 desaturases.
Abbreviations are: Rcfah12, fah12 hydroxylase gene
from R. communis (van de Loo et al., 1995); Lffah12,
kappa hydroxylase gene from L. fendleri; Atfad2,
fad2 desaturase from Arabidopsis thaliana (Okuley et

- al., 1994); Gmfad2-1, fad2 desaturase from Glycine max (GenBank accession number L43920); Gmfad2-2, fad2 desaturase from Glycine max (Genbank accession number L43921); Zmfad2, fad2 desaturase from Zea
- mays (PCT WO 94/11516); Rcfad2, fragment of fad2 desaturase from R. communis (PCT WO 94/11516); Bnfad2, fad2 desaturase from Brassica napus (PCT WO 94/11516); LFFAH12.AMI, SEQ ID NO:4; FAH12.AMI, SEQ ID NO:5; ATFAD2.AMI, SEQ ID NO:6; BNFAD2.AMI, SEQ ID
- NO:7; GMFAD2-1.AMI, SEQ ID NO:8; GMFAD2-2.AMI, SEQ ID NO:9; ZMFAD2.AMI, SEQ ID NO:10; and RCFAD2.AMI, SEQ ID NO:11.

Figure 10 shows a Southern blot of genomic DNA from L. fendleri probed with pLesq-HYD. E = EcoRI, H = HindIII, X = XbaI.

Figure 11 shows a map of binary Ti plasmid pSLJ44024.

Figure 12 shows a map of plasmid pYES2.0

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Figure 13 shows part of a gas chromatogram of derivatized fatty acids from yeast cells that contain plasmid plesqYes in which expression of the hydroxylase gene was induced by addition of galactose to the growth medium. The arrow points to a peak that is not present in uninduced cells. The lower part of the figure is the mass spectrum of the peak indicated by the arrow.

SUMMARY OF THE INVENTION

10 This invention relates to plant fatty acyl hydroxylases. Methods to use conserved amino acid or nucleotide sequences to obtain plant fatty acyl hydroxylases are described. Also described is the use of cDNA clones encoding a plant hydroxylase to produce a family of hydroxylated fatty acids in transgenic plants.

In a first embodiment, this invention is directed to recombinant DNA constructs which can provide for the transcription, or transcription and translation (expression) of the plant kappa hydroxylase sequence. In particular, constructs which are capable of transcription, or transcription and translation in plant host cells are preferred. Such constructs may contain a variety of regulatory regions including transcriptional initiation regions obtained from genes preferentially expressed in plant seed tissue. In a second aspect, this invention relates to the presence of such constructs in host cells, especially plant host cells which have an expressed plant kappa hydroxylase therein.

In yet another aspect, this invention relates to a method for producing a plant kappa hydroxylase in a host cell or progeny thereof via the expression

of a construct in the cell. Cells containing a plant kappa hydroxylase as a result of the production of the plant kappa hydroxylase encoding sequence are also contemplated herein.

In another embodiment, this invention relates to methods of using a DNA sequence encoding a plant kappa hydroxylase for the modification of the proportion of hydroxylated fatty acids produced within a cell, especially plant cells. Plant cells having such a modified hydroxylated fatty acid composition are also contemplated herein.

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In a further aspect of this invention, plant kappa hydroxylase proteins and sequences which are related thereto, including amino acid and nucleic acid sequences, are contemplated. Plant kappa hydroxylase exemplified herein includes a Lesquerella fendleri fatty acid hydroxylase. This exemplified fatty acid hydroxylase may be used to obtain other plant fatty acid hydroxylases of this invention.

In a further aspect of this invention, a nucleic acid sequence which directs the seed specific expression of an associated polypeptide coding sequence is described. The use of this nucleic acid sequence or fragments derived therefrom, to obtain seed-specific expression in higher plants of any coding sequence is contemplated herein.

In a further aspect of this invention, the

use of genes encoding fatty acyl hydroxylases of
this invention are used to alter the amount of fatty
acid unsaturation of seed lipids. The present
invention further discloses the use of genetically
modified hydroxylase and desaturase genes to achieve

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directed modification of fatty acid unsaturation levels.

DETAILED DESCRIPTION OF THE INVENTION

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to function.

A genetically transformed plant of the present invention which accumulates hydroxylated fatty acids can be obtained by expressing the double-stranded DNA molecules described in this application.

A plant fatty acid hydroxylase of this 10 invention includes any sequence of amino acids, such as a protein, polypeptide or peptide fragment, or nucleic acid sequences encoding such polypeptides, obtainable from a plant source which demonstrates the ability to catalyze the production of ricinoleic, lesquerolic, hydroxyerucic (16-15 hydroxydocos-cis-13-enoic acid) or hydroxypalmitoleic (12-hydroxyhexadec-cis-9-enoic) from CoA, ACP or lipid-linked monoenoic fatty acid substrates under plant enzyme reactive conditions. 20 By "enzyme reactive conditions" is meant that any necessary conditions are available in an environment (i.e., such factors as temperature, pH, lack of

Preferential activity of a plant fatty acid hydroxylase toward a particular fatty acyl substrate is determined upon comparison of hydroxylated fatty acid product amounts obtained per different fatty acyl substrates. For example, by "oleate preferring" is meant that the hydroxylase activity of the enzyme preparation demonstrates a preference for oleate-containing substrates over other substrates. Although the precise substrate of the castor fatty

inhibiting substances) which will permit the enzyme

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acid hydroxylase is not known, it is thought to be a monounsaturated fatty acid moiety which is esterified to a phospholipid such as phosphatidylcholine. However, it is also possible that monounsaturated fatty acids esterified to phosphatidylethanolamine, phosphatidic acid or a neutral lipid such as diacylglycerol or a Coenzyme-A thioester may also be substrates.

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As noted above, significant activity has been 10 observed in radioactive labelling studies using fatty acyl substrates other than oleate (Howling et al., 1972) indicating that the substrate specificity is for a family of related fatty acyl compounds. Because the castor hydroxylase introduces hydroxy 15 groups three carbons from a double bond, proximal to the methyl carbon of the fatty acid, the enzyme is termed a kappa hydroxylase for convenience. Of particular interest, the present invention discloses that the castor kappa hydroxylase may be used for 20 production of 12-hydroxy-9-octadecenoic acid (ricinoleate), 12-hydroxy-9-hexadecenoic acid, 14hydroxy-11-eicosenoic acid, 16-hydroxy-13-docosenoic acid, 9-hydroxy-6-octadecenoic acid by expression in plants species which produce the non-hydroxylated 25 precursors. The present invention also discloses production of additionally modified fatty acids such as 12-hydroxy-9,15-octadecadienoic acid that result from desaturation of hydroxylated fatty acids (e.g., 12-hydroxy-9-octadecenoic acid in this example).

The present invention also discloses that future advances in the genetic engineering of plants will lead to production of substrate fatty acids, such as icosenoic acid esters, and palmitoleic acid esters in plants that do not normally accumulate

such fatty acids. The invention described herein may be used in conjunction with such future improvements to produce hydroxylated fatty acids of this invention in any plant species that is amenable to directed genetic modification. Thus, the applicability of this invention is not limited in our conception only to those species that currently accumulate suitable substrates.

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As noted above, a plant kappa hydroxylase of 10 this invention will display activity towards various fatty acyl substrates. During biosynthesis of lipids in a plant cell, fatty acids are typically covalently bound to acyl carrier protein (ACP), coenzyme A (CoA) or various cellular lipids. Plant kappa hydroxylases which display preferential 15 activity toward lipid-linked acyl substrate are especially preferred because they are likely to be closely associated with normal pathway of storage lipid synthesis in immature embryos. However, activity toward acyl-CoA substrates or other 20 synthetic substrates, for example, is also contemplated herein.

Other plant kappa hydroxylases are obtainable from the specific exemplified sequences provided herein. Furthermore, it will be apparent that one can obtain natural and synthetic plant kappa hydroxylases including modified amino acid sequences and starting materials for synthetic-protein modeling from the exemplified plant kappa hydroxylase and from plant kappa hydroxylases which are obtained through the use of such exemplified sequences. Mcdified amino acid sequences include sequences which have been mutated, truncated, elongated or the like, whether such sequences were

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partially or wholly synthesized. Sequences which are actually purified from plant preparations or are identical or encode identical proteins thereto, regardless of the method used to obtain the protein or sequence, are equally considered naturally derived.

Thus, one skilled in the art will readily recognize that antibody preparations, nucleic acid probes (DNA and RNA) or the like may be prepared and 10 used to screen and recover "homologous" or "related" kappa hydroxylases from a variety of plant sources. Typically, nucleic acid probes are labeled to allow detection, preferably with radioactivity although enzymes or other methods may also be used. For 15 immunological screening methods, antibody preparations either monoclonal or polyclonal are utilized. Polyclonal antibodies, although less specific, typically are more useful in gene isolation. For detection, the antibody is labeled using radioactivity or any one of a variety of 20 second antibody/enzyme conjugate systems that are commercially available.

Homologous sequences are found when there is an identity of sequence and may be determined upon comparison of sequence information, nucleic acid or amino acid, or through hybridization reactions between a known kappa hydroxylase and a candidate source. Conservative changes, such as Glu/Asp, Val/Ile, Ser/Thr, Arg/Lys and Gln/Asn may also be considered in determining sequence homology.

Typically, a lengthy nucleic acid sequence may show as little as 50-60% sequence identity, and more preferably at least about 70% sequence identity, between the target sequence and the given plant

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kappa hydroxylase of interest excluding any deletions which may be present, and still be considered related. Amino acid sequences are considered homologous by as little as 25% sequence identity between the two complete mature proteins. (see generally, Doolittle, R.F., OF URFS and ORFS, University Science Books, CA, 1986.)

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A genomic or other appropriate library prepared from the candidate plant source of interest 10 may be probed with conserved sequences from the plant kappa hydroxylase to identify homologously related sequences. Use of an entire cDNA or other sequence may be employed if shorter probe sequences are not identified. Positive clones are then 15 analyzed by restriction enzyme digestion and/or sequencing. When a genomic library is used, one or more sequences may be identified providing both the coding region, as well as the transcriptional regulatory elements of the kappa hydroxylase gene 20 from such plant source. Probes can also be considerably shorter than the entire sequence. Oligonucleotides may be used, for example, but should be at least about 10, preferably at least about 15, more preferably at least 20 nucleotides in 25 length. When shorter length regions are used for comparison, a higher degree of sequence identity is required than for longer sequences. Shorter probes are often particularly useful for polymerase chain reactions (PCR), especially when highly conserved 30 sequences can be identified (see Gould et al., 1989 for examples of the use of PCR to isolate homologous genes from taxonomically diverse species).

When longer nucleic acid fragments are employed (>100 bp) as probes, especially when using

complete or large cDNA sequences, one would screen with low stringencies (for example, 40-50°C below the melting temperature of the probe) in order to obtain signal from the target sample with 20-50% deviation, i.e., homologous sequences (Beltz et al., 1983).

In a preferred embodiment, a plant kappa hydroxylase of this invention will have at least 60% overall amino acid sequence similarity with the exemplified plant kappa hydroxylase. In particular, 10 kappa hydroxylases which are obtainable from an amino acid or nucleic acid sequence of a castor or Lesquerella kappa hydroxylase are especially preferred. The plant kappa hydroxylases may have preferential activity toward longer or shorter chain 15 fatty acyl substrates. Plant fatty acyl hydroxylases having oleate-12-hydroxylase activity and eicosenoate-14-hydroxylase activity are both considered homologously related proteins because of in vitro evidence (Howling et al., 1972), and 20 evidence disclosed herein, that the castor kappa hydroxylase will act on both substrates. Hydroxylated fatty acids may be subject to further enzymatic modification by other enzymes which are normally present or are introduced by genetic 25 engineering methods. For example, 14-hydroxy-11,17eicosadienoic acid, which is present in some Lesquerella species (Smith, 1985), is thought to be produced by desaturation of 14-hydroxy-11-eicosenoic 30 acid.

Again, not only can gene clones and materials derived therefrom be used to identify homologous plant fatty acyl hydroxylases, but the resulting sequences obtained therefrom may also provide a

hydroxylases from other plant sources. In particular, PCR may be a useful technique to obtain related plant fatty acyl hydroxylases from sequence data provided herein. One skilled in the art will be able to design oligonucleotide probes based upon sequence comparisons or regions of typically highly conserved sequence. Of special interest are polymerase chain reaction primers based on the conserved regions of amino acid sequence between the castor kappa hydroxylase and the *L. fendleri* hydroxylase (SEQ ID NO:4). Details relating to the design and methods for a PCR reaction using these probes are described more fully in the examples.

It should also be noted that the fatty acyl hydroxylases of a variety of sources can be used to investigate fatty acid hydroxylation events in a wide variety of plant and in vivo applications. Because all plants synthesize fatty acids via a common metabolic pathway, the study and/or application of one plant fatty acid hydroxylase to a heterologous plant host may be readily achieved in a variety of species.

Once the nucleic acid sequence is obtained, the transcription, or transcription and translation (expression), of the plant fatty acyl hydroxylases in a host cell is desired to produce a ready source of the enzyme and/or modify the composition of fatty acids found therein in the form of free fatty acids, esters (particularly esterified to glycerolipids or as components of wax esters), estolides, or ethers. Other useful applications may be found when the host cell is a plant host cell, in vitro and in vivo. For example, by increasing the amount of an kappa

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hydroxylase available to the plant, an increased percentage of ricinoleate or lesqueroleate (14-hydroxy-11-eicosenoic acid) may be provided.

Kappa Hydroxylase

5 By this invention, a mechanism for the biosynthesis of ricinoleic acid in plants is demonstrated. Namely, that a specific plant kappa hydroxylase having preferential activity toward fatty acyl substrates is involved in the accumulation of hydroxylated fatty acids in at least 10 some plant species. The use of the terms ricinoleate or ricinoleic acid (or lesqueroleate or lesquerolic acid, densipoleate etc.) is intended to include the free acids, the ACP and CoA esters, the salts of 15 these acids, the glycerolipid esters (particularly the triacylglycerol esters), the wax esters, the estolides and the ether derivatives of these acids.

The determination that plant fatty acyl hydroxylases are active in the *in vivo* production of hydroxylated fatty acids suggests several possibilities for plant enzyme sources. In fact, hydroxylated fatty acids are found in some natural plant species in abundance. For example, three hydroxy fatty acids related to ricinoleate occur in major amounts in seed oils from various *Lesquerella* species. Of particular interest, lesquerolic acid is a 20 carbon homolog of ricinoleate with two additional carbons at the carboxyl end of the chain (Smith, 1985). Other natural plant sources of hydroxylated fatty acids include but are not limited to seeds of the *Linum* genus, seeds of *Wrightia* species, *Lycopodium* species, *Strophanthus* species,

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Convolvulaces species, Calendula species and many others (van de Loo et al., 1993).

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Plants having significant presence of ricinoleate or lesqueroleate or desaturated other or modified derivatives of these fatty acids are preferred candidates to obtain naturally-derived kappa hydroxylases. For example, Lesquerella densipila contains a diunsaturated 18 carbon fatty acid with a hydroxyl group (van de Loo et al., 1993) that is thought to be produced by an enzyme that is closely related to the castor kappa hydroxylase, according to the theory on which this invention is based. In addition, a comparison between kappa hydroxylases and between plant fatty acvl hydroxylases which introduce hydroxyl groups at positions other than the 12-carbon of oleate or the 14-carbon of lesqueroleate or on substrates other than oleic acid and icosenoic acid may yield insights for gene identification, protein modeling or other modifications as discussed above.

Especially of interest are fatty acyl hydroxylases which demonstrate activity toward fatty acyl substrates other than oleate, or which introduce the hydroxyl group at a location other than the C12 carbon. As described above, other plant sources may also provide sources for these enzymes through the use of protein purification, nucleic acid probes, antibody preparations, protein modeling, or sequence comparisons, for example, and of special interest are the respective amino acid and nucleic acid sequences corresponding to such plant fatty acyl hydroxylases. Also, as previously described, once a nucleic acid sequence is obtained for the given plant hydroxylase, further plant

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sequences may be compared and/or probed to obtain homologously related DNA sequences thereto and so on.

Genetic Engineering Applications

WO 97/30582

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As is well known in the art, once a cDNA clone encoding a plant kappa hydroxylase is obtained, it may be used to obtain its corresponding genomic nucleic acid sequences thereto.

The nucleic acid sequences which encode plant 10 kappa hydroxylases may be used in various constructs, for example, as probes to obtain further sequences from the same or other species. Alternatively, these sequences may be used in conjunction with appropriate regulatory sequences to increase levels of the respective hydroxylase of 15 interest in a host cell for the production of hydroxylated fatty acids or study of the enzyme in vitro or in vivo or to decrease or increase levels of the respective hydroxylase of interest for some 20 applications when the host cell is a plant entity, including plant cells, plant parts (including but not limited to seeds, cuttings or tissues) and plants.

A nucleic acid sequence encoding a plant kappa hydroxylase of this invention may include genomic, cDNA or mRNA sequence. By "encoding" is meant that the sequence corresponds to a particular amino acid sequence either in a sense or anti-sense orientation. By "recombinant" is meant that the sequence contains a genetically engineered modification through manipulation via mutagenesis, restriction enzymes, or the like. A cDNA sequence may or may not encode pre-processing sequences, such

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as transit or signal peptide sequences. Transit or signal peptide sequences facilitate the delivery of the protein to a given organelle and are frequently cleaved from the polypeptide upon entry into the organelle, releasing the "mature" sequence. The use of the precursor DNA sequence is preferred in plant cell expression cassettes.

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Furthermore, as discussed above the complete genomic sequence of the plant kappa hydroxylase may be obtained by the screening of a genomic library with a probe, such as a cDNA probe, and isolating those sequences which regulate expression in seed tissue.

Once the desired plant kappa hydroxylase 15 nucleic acid sequence is obtained, it may be manipulated in a variety of ways. Where the sequence involves non-coding flanking regions, the flanking regions may be subjected to resection, mutagenesis, etc. Thus, transitions, transversions, deletions, 20 and insertions may be performed on the naturally occurring sequence. In addition, all or part of the sequence may be synthesized. In the structural gene, one or more codons may be modified to provide for a modified amino acid sequence, or one or more codon 25 mutations may be introduced to provide for a convenient restriction site or other purpose involved with construction or expression. The structural gene may be further modified by employing synthetic adapters, linkers to introduce one or more convenient restriction sites, or the like. 30

The nucleic acid or amino acid sequences encoding a plant kappa hydroxylase of this invention may be combined with other non-native, or "heterologous", sequences in a variety of ways. By

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"heterologous" sequences is meant any sequence which is not naturally found joined to the plant kappa hydroxylase, including, for example, combination of nucleic acid sequences from the same plant which are not naturally found joined together.

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The DNA sequence encoding a plant kappa hydroxylase of this invention may be employed in conjunction with all or part of the gene sequences normally associated with the kappa hydroxylase. In its component parts, a DNA sequence encoding kappa hydroxylase is combined in a DNA construct having, in the 5' to 3' direction of transcription, a transcription initiation control region capable of promoting transcription and/or translation in a host cell, the DNA sequence encoding plant kappa hydroxylase and a transcription and/or translation termination region.

Potential host cells include both prokaryotic and eukaryotic cells. A host cell may be unicellular or found in a multicellular differentiated or undifferentiated organism depending upon the intended use. Cells of this invention may be distinguished by having a plant kappa hydroxylase foreign to the wild-type cell present therein, for example, by having a recombinant nucleic acid construct encoding a plant kappa hydroxylase therein.

Depending upon the host, the regulatory regions will vary, including regions from viral,

30 plasmid or chromosomal genes, or the like. For expression in prokaryotic or eukaryotic microorganisms, particularly unicellular hosts, a wide variety of constitutive or regulatable promoters may be employed. Expression in a

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microorganism can provide a ready source of the plant enzyme. Among transcriptional initiation regions which have been described are regions from bacterial and yeast hosts, such as E. coli, B. subtilis, Saccharomyces cerevisiae, including genes such as beta-galactosidase, T7 polymerase, trpE or the like.

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involve regulatory regions functional in plants
which provide for modified production of plant kappa
hydroxylase with resulting modification of the fatty
acid composition. The open reading frame, coding for
the plant kappa hydroxylase or functional fragment
thereof will be joined at its 5' end to a
transcription initiation regulatory region. Numerous
transcription initiation regions are available which
provide for a wide variety of constitutive or
regulatable, e.g., inducible, transcription of the
structural gene functions.

Among transcriptional initiation regions used for plants are such regions associated with the structural genes such as for nopaline and mannopine synthases, or with napin, soybean β -conglycinin, oleosin, 12S storage protein, the cauliflower mosaic virus 35S promoters or the like. The transcription/translation initiation regions corresponding to such structural genes are found immediately 5' upstream to the respective start codons.

In embodiments wherein the expression of the kappa hydroxylase protein is desired in a plant host, the use of all or part of the complete plant kappa hydroxylase gene is desired. If a different promoter is desired, such as a promoter native to the plant host of interest or a modified promoter,

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i.e., having transcription initiation regions derived from one gene source and translation initiation regions derived from a different gene source or enhanced promoters, such as double 35S CaMV promoters, the sequences may be joined together using standard techniques.

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For such applications when 5' upstream noncoding regions are obtained from other genes
regulated during seed maturation, those

10 preferentially expressed in plant embryo tissue,
such as transcription initiation control regions
from the B. napus napin gene, or the Arabidopsis 12S
storage protein, or soybean β-conglycinin (Bray et
al., 1987) are desired. Transcription initiation

15 regions which are preferentially expressed in seed
tissue, i.e., which are undetectable in other plant
parts, are considered desirable for fatty acid
modifications in order to minimize any disruptive or
adverse effects of the gene product.

20 Regulatory transcript termination regions may be provided in DNA constructs of this invention as well. Transcript termination regions may be provided by the DNA sequence encoding the plant kappa hydroxylase or a convenient transcriptiontermination region derived from a different gene 25 source, for example, the transcript termination region which is naturally associated with the transcript initiation region. Where the transcript termination region is from a different gene source, it will contain at least about 0.5 kb, preferably 30 about 1-3 kb of sequence 3' to the structural gene from which the termination region is derived.

Plant expression or transcription constructs having a plant kappa hydroxylase as the DNA sequence

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of interest for increased or decreased expression thereof may be employed with a wide variety of plant life, particularly, plant life involved in the production of vegetable oils for edible and industrial uses. Most especially preferred are temperate oilseed crops. Plants of interest include, but are not limited to rapeseed (canola and high erucic acid varieties), Crambe, Brassica juncea, Brassica nigra, meadowfoam, flax, sunflower, safflower, cotton, Cuphea, soybean, peanut, coconut and oil palms and corn. An important criterion in the selection of suitable plants for the introduction on the kappa hydroxylase is the presence in the host plant of a suitable substrate for the hydroxylase. Thus, for example, production of ricinoleic acid will be best accomplished in plants that normally have high levels of oleic acid

Depending on the method for introducing the recombinant constructs into the host cell, other DNA sequences may be required. Importantly, this invention is applicable to dicotyledons and monocotyledons species alike and will be readily applicable to new and/or improved transformation and regulation techniques. The method of transformation is not critical to the current invention; various methods of plant transformation are currently available. As newer methods are available to transform crops, they may be directly applied hereunder. For example, many plant species naturally susceptible to Agrobacterium infection may be successfully transformed via tripartite or binary

in seed lipids. Similarly, production of lesquerolic acid will best be accomplished in plants that have

high levels of icosenoic acid in seed lipids.

vector methods of Agrobacterium mediated transformation. In addition, techniques of microinjection, DNA particle bombardment, electroporation have been developed which allow for the transformation of various monocot and dicot plant species.

In developing the DNA construct, the various components of the construct or fragments thereof will normally be inserted into a convenient cloning vector which is capable of replication in a bacterial host, e.g., E. coli. Numerous vectors exist that have been described in the literature. After each cloning, the plasmid may be isolated and subjected to further manipulation, such as restriction, insertion of new fragments, ligation, deletion, insertion, resection, etc., so as to tailor the components of the desired sequence. Once the construct has been completed, it may then be transferred to an appropriate vector for further manipulation in accordance with the manner of transformation of the host cell.

Normally, included with the DNA construct will be a structural gene having the necessary regulatory regions for expression in a host and providing for selection of transformant cells. The gene may provide for resistance to a cytotoxic agent, e.g., antibiotic, heavy metal, toxin, etc., complementation providing prototropy to an auxotrophic host, viral immunity or the like.

Depending upon the number of different host species the expression construct or components thereof are introduced, one or more markers may be employed, where different conditions for selection are used for the different hosts.

It is noted that the degeneracy of the DNA code provides that some codon substitutions are permissible of DNA sequences without any corresponding modification of the amino acid sequence.

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As mentioned above, the manner in which the DNA construct is introduced into the plant host is not critical to this invention. Any method which provides for efficient transformation may be employed. Various methods for plant cell 10 transformation include the use of Ti- or Riplasmids, microinjection, electroporation, infiltration, imbibition, DNA particle bombardment, liposome fusion, DNA bombardment or the like. In many instances, it will be desirable to have the 15 construct bordered on one or both sides of the T-DNA, particularly having the left and right borders, more particularly the right border. This is particularly useful when the construct uses A. tumefaciens or A. rhizogenes as a mode for 20 transformation, although the T-DNA borders may find use with other modes of transformation.

Where Agrobacterium is used for plant cell transformation, a vector may be used which may be introduced into the Agrobacterium host for homologous recombination with T-DNA or the Ti- or Ri-plasmid present in the Agrobacterium host. The Ti- or Ri-plasmid containing the T-DNA for recombination may be armed (capable of causing gall formation) or disarmed (incapable of causing gall), the latter being permissible, so long as the vir genes are present in the transformed Agrobacterium host. The armed plasmid can give a mixture of normal plant cells and gall.

In some instances where Agrobacterium is used as the vehicle for transforming plant cells, the expression construct bordered by the T-DNA border(s) will be inserted into a broad host spectrum vector. there being broad host spectrum vectors described in 5 the literature. Commonly used is pRK2 or derivatives thereof. See, for example, Ditta et al. (1980), which is incorporated herein by reference. Included with the expression construct and the T-DNA will be one or more markers, which allow for selection of 10 transformed Agrobacterium and transformed plant cells. A number of markers have been developed for use with plant cells, such as resistance to kanamycin, the aminoglycoside G418, hygromycin, or the like. The particular marker employed is not 15 essential to this invention, one or another marker being preferred depending on the particular host and the manner of construction.

For transformation of plant cells using Agrobacterium, explants may be combined and 20 incubated with the transformed Agrobacterium for sufficient time for transformation, the bacteria killed, and the plant cells cultured in an appropriate selective medium. Once callus forms, 25 shoot formation can be encouraged by employing the appropriate plant hormones in accordance with known methods and the shoots transferred to rooting medium for regeneration of plants. The plants may then be grown to seed and the seed used to establish 30 repetitive generations and for isolation of vegetable oils.

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Using Hydroxylase Genes to Alter the Activity of Fatty Acid Desaturases

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A widely acknowledged goal of current efforts to improve the nutritional quality of edible plant oils, or to facilitate industrial applications of plant oils, is to alter the level of desaturation of plant storage lipids (Topfer et al., 1995). In particular, in many crop species it is considered desirable to reduce the level of polyunsaturation of storage lipids and to increase the level of oleic acid. The precise amount of the various fatty acids in a particular plant oil varies with the intended application. Thus, it is desirable to have a robust method that will permit genetic manipulation of the level of unsaturation to any desired level.

Substantial progress has recently been made in the isolation of genes encoding plant fatty acid desaturases (reviewed in Topfer et al., 1995). These genes have been introduced into various plant species and used to alter the level of fatty acid unsaturation in one of three ways. First, the genes can be placed under transcriptional control of a strong promoter so that the amount of the corresponding enzyme is increased. In some cases this leads to an increase in the amount of the fatty acid that is the product of the reaction catalyzed by the enzyme. For example, Arondel et al. (1992) increased the amount of linolenic acid (18:3) in tissues of transgenic Arabidopsis plants by placing the endoplasmic reticulum-localized fad3 gene under transcriptional control of the strong constitutive cauliflower mosaic virus 35S promoter.

A second method of using cloned genes to alter the level of fatty acid unsaturation is to

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cause transcription of all or part of a gene in transgenic tissues so that the transcripts have an antisense orientation relative to the normal mode of transcription. This has been used by a number of laboratories to reduce the level of expression of one or more desaturase genes that have significant nucleotide sequence homology to the gene used in the construction of the antisense gene (reviewed in Topfer et al.). For instance, antisense repression of the oleate $\Delta 12$ -desaturase in transgenic rapeseed resulted in a strong increase in oleic acid content (cf., Topfer et al., 1995).

A third method for using cloned genes to alter fatty acid desaturation is to exploit the phenomenon of cosuppression or "gene-silencing" (Matzke et al., 1995). Although the mechanisms responsible for gene silencing are not known in any detail, it has frequently been observed that in transgenic plants, expression of an introduced gene leads to inactivation of homologous endogenous genes.

For example, high-level sense expression of the Arabidopsis fad8 gene, which encodes a chloroplast-localized $\Delta 15$ -desaturase, in transgenic Arabidopsis plants caused suppression of the endogenous copy of the fad8 gene and the homologous fad7 gene (which encodes an isozyme of the fad8 gene) (Gibson et al., 1994). The fad7 and fad8 genes are only 76% identical at the nucleotide level. At the time of publication, this example represented the most divergent pair of plant genes for which cosuppression had been observed.

In view of previous evidence concerning the relatively high level of nucleotide sequence

homology required to obtain cosuppression, it is not obvious to one skilled in the art that sense expression in transgenic plants of the castor fatty acyl hydroxylase of this invention would significantly alter the amount of unsaturation of storage lipids.

However, the present inventors establish that fatty acyl hydroxylase genes can be used for this purpose as taught in Example 4 of this specification. Of particular importance, this invention teaches the use of fatty acyl hydroxylase genes to increase the proportion of oleic acid in transgenic plant tissues. The mechanism by which expression of the gene exerts this effect is not known but may be due to one of several possibilities

The invention now being generally described, it will be more readily understood by reference to the following examples which are included for purposes of illustration only and are not intended to limit the present invention.

which are elaborated upon in Example 4.

EXAMPLES

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In the experimental disclosure which follows, all temperatures are given in degrees centigrade (°C), weights are given in grams (g), milligram (mg) or micrograms (μ g), concentrations are given as molar (M), millimolar (mM) or micromolar (μ M) and all volumes are given in liters (l), microliters (μ l) or milliliters (ml), unless otherwise indicated.

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EXAMPLE 1 - PRODUCTION OF NOVEL HYDROXYLATED FATTY ACIDS IN ARABIDOPSIS THALIANA Overview

The kappa hydroxylase encoded by the fah12 gene from castor was used to produce ricinoleic acid, lesquerolic acid, densipolic acid and auricolic acid in transgenic Arabidopsis plants.

Production of transgenic plants

A variety of methods have been developed to

insert a DNA sequence of interest into the genome of
a plant host to obtain the transcription and
translation of the sequence to effect phenotypic
changes. The following methods represent only one of
many equivalent means of producing transgenic plants
and causing expression of the hydroxylase gene.

Arabidopsis plants were transformed, by Agrobacterium-mediated transformation, with the kappa hydroxylase encoded by the castor fahl2 gene on binary Ti plasmid pB6. This plasmid has also been used to transform Nicotiana tabacum for the production of ricinoleic acid.

Inoculums of Agrobacterium tumefaciens strain GV3101 containing binary Ti plasmid pB6 were plated on L-broth plates containing 50 µg/ml kanamycin and incubated for 2 days at 30°C. Single colonies were used to inoculate large liquid cultures (L-broth medium with 50 mg/l rifampicin, 110 mg/l gentamycin and 200 mg/l kanamycin) to be used for the transformation of Arabidopsis plants.

Arabidopsis plants were transformed by the in planta transformation procedure essentially as described by Bechtold et al. (1993). Cells of A. tumefaciens GV3101(pB6) were harvested from liquid

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cultures by centrifugation, then resuspended in infiltration medium at $OD_{600} = 0.8$. Infiltration medium was Murashige and Skoog macro and micronutrient medium (Sigma Chemical Co., St. Louis, 5 MO) containing 10 mg/l 6-benzylaminopurine and 5% glucose. Batches of 12-15 plants were grown for 3 to 4 weeks in natural light at a mean daily temperature of approximately 25°C in 3.5 inch pots containing soil. The intact plants were immersed in the bacterial suspension then transferred to a vacuum 10 chamber and placed under 600 mm of vacuum produced by a laboratory vacuum pump until tissues appeared uniformly water-soaked (approximately 10 min). The plants were grown at 25°C under continuous light (100 μ mol m⁻² s⁻¹ irradiation in the 400 to 700 nm 15 range) for four weeks. The seeds obtained from all the plants in a pot were harvested as one batch. The seeds were sterilized by sequential treatment for 2 min with ethanol followed by 10 min in a mixture of household bleach (Chlorox), water and Tween-80 (50%, 20 50%, 0.05%) then rinsed thoroughly with sterile water. The seeds were plated at high density (2000 to 4000 per plate) onto agar-solidified medium in 100 mm petri plates containing 1/2 X Murashige and Skoog salts medium enriched with B5 vitamins (Sigma 25 Chemical Co., St. Louis, MO) and containing kanamycin at 50 mg/l. After incubation for 48 h at 4°C to stimulate germination, seedlings were grown for a period of seven days until transformants were clearly identifiable as healthy green seedlings 30 against a background of chlorotic kanamycinsensitive seedlings. The transformants were transferred to soil for two weeks before leaf tissue could be used for DNA and lipid analysis. More than 20 transformants were obtained.

DNA was extracted from young leaves from transformants to verify the presence of an intact fah12 gene. The presence of the transgene in a 5 number of the putative transgenic lines was verified by using the polymerase chain reaction to amplify the insert from pB6. The primers used were HF2 = GCTCTTTTGTGCGCTCATTC (SEQ ID NO:12) and HR1 = 10 CGGTACCAGAAAACGCCTTG (SEQ ID NO:13), which were designed to allow the amplification of a 700 bp fragment. Approximately 100 ng of genomic DNA was added to a solution containing 25 pmol of each primer, 1.5 U Taq polymerase (Boehringer Manheim), 200 uM of dNTPs, 50 mM KCl, 10 mM Tris.Cl (pH 9), 15 0.1% (v/v) Triton X-100, 1.5 mM MgCl₂, 3% (v/v)formamide, to a final volume of 50 μ l. Amplifications conditions were: 4 min denaturation step at 94°C, followed by 30 cycles of 92°C for 1 min, 55°C for 1 min, 72°C for 2 min. A final 20 extension step closed the program at 72°C for 5 min. Transformants could be positively identified after visualization of a characteristic 1 kb amplified fragment on an ethidium bromide stained agarose gel. All transgenic lines tested gave a PCR product of a 25 size consistent with the expected genotype, confirming that the lines were, indeed, transgenic: All further experiments were done with three representative transgenic lines of the wild type 30 designated as 1-3, 4D, 7-4 and one transgenic line of the fad2 mutant line JB12. The transgenic JB12 line was included in order to test whether the increased accumulation of oleic acid in this mutant

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would have an effect on the amount of ricinoleic acid that accumulated in the transgenic plants.

Analysis of transgenic plants

Leaves and seeds from fahl2 transgenic Arabidopsis plants were analyzed for the presence of 5 hydroxylated fatty acids using gas chromatography. Lipids were extracted from 100-200 mg leaf tissue or 50 seeds. Fatty acid methyl esters (FAMES) were prepared by placing tissue in 1.5 ml of 1.0 M 10 methanolic HCl (Supelco Co.) in a 13 x 100 mm glass screw-cap tube capped with a teflon-lined cap and heated to 80°C for 2 hours. Upon cooling, 1 ml petroleum ether was added and the FAMES removed by aspirating off the ether phase which was then dried under a nitrogen stream in a glass tube. One hundred 15 μl of N,O-bis(Trimethylsilyl) trifluoroacetamide (BSTFA; Pierce Chemical Co) and 200 μ l acetonitrile was added to derivatize the hydroxyl groups. The reaction was carried out at 70°C for 15 min. The products were dried under nitrogen, redissolved in 20 100 µl chloroform and transferred to a gas chromatograph vial. Two μ l of each sample were analyzed on a SP2340 fused silica capillary column (30 m, 0.75 mm ID, 0.20 mm film, Supelco), using a Hewlett-Packard 5890 II series Gas Chromatograph. 25 The samples were not split, the temperature program was 195°C for 18 min, increased to 230°C at 25°C/min, held at 230°C for 5 min then down to 195°C at 25°C/min., and flame ionization detectors were 30 used.

The chromatographic elution time of methyl esters and O-TMS derivatives of ricinoleic acid, lesquerolic acid and suricolic acid was established

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by GC-MS of lipid samples from seeds of *L. fendleri* and comparison to published chromatograms of fatty acids from this species (Carlson et al., 1990). A O-TMS-methyl-ricinoleate standard was prepared from ricinoleic acid obtained from Sigma Chemical Co (St, Louis, MO). O-TMS-methyl-lesqueroleate and O-TMS-methyl-auricoleate standards were prepared from triacylglycerols purified from seeds of *L. fendleri*. The mass spectrum of O-TMS-methyl-ricinoleate, O-TMS-methyl-densipoleate, O-TMS-methyl-lesqueroleate, and O-TMS-methyl-auricoleate are shown in Figures 1A-D, respectively. The structures of the characteristic ions produced during mass spectrometry of these derivatives are shown in Figure 2.

Lipid extracted from transgenic tissues were analyzed by gas chromatography and mass spectrometry for the presence of hydroxylated fatty acids. As a matter of reference, the average fatty acid 20 composition of leaves in Arabidopsis wild type and fad2 mutant lines was reported by Miguel and Browse (1992). Gas chromatograms of methylated and silylated fatty acids from seeds of wild type and a fah12 transgenic wild type plant are shown in Figures 3A and 3B, respectively. The profiles are 25 very similar except for the presence of three small but distinct peaks at 14.3, 15.9 and 18.9 minutes. A very small peak at 20.15 min was also evident. The elution time of the peaks at 14.3 and 18.9 min corresponded precisely to that of comparably 30 prepared ricinoleic and lesquerolic standards, respectively. No significant differences were observed in lipid extracts from leaves or roots of

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the wild type and the fahl2 transgenic wild type lines (Table 1).

Thus, in spite of the fact that the fahl2 gene is expressed throughout the plant, effects on fatty acid composition was observed only in seed tissue. The present inventors have made a similar observation for transgenic fahl2 tobacco.

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Table 1. Fatty acid composition of lipids from transgenic and wild type Arabidopsis. The values are the means obtained from analysis of samples from three independent transgenic lines, or three independent samples of wild type and fad2 lines.

TABLE 1

	Root	FAH12 WT	24.9	0	1.9	3.2	29.4	
		WT	23.9	0	2.0	5.4	32.2	
	Leaf	FAII12 W.F	17.5	9.8	1.2	3.4	14.0	
	Le	WT	16.5	10.1	1.3	2.4	15.1	
	Seed	JB12	6.1	0	3.5	47.8	7.2	
		FAH12 fad2	6.4	0	2.9	43.4	10.2	
		FAH12 WT	8.2	0	3.5	26.3	21.4	
	:	IM	8.5	0	3.2	15.4	27.0	
Fatty	acid		16:0	16:3	18:0	18:1	18:2	

TABLE 1 (continued)

Root	FAH12 WT	0	0	0	0
, a	ΤM	0	0	0	0
Leaf	FAH12 WT	0	0	0	0
J	LМ	0	0	0	0
	JB12	0	0	0	0
Seed	FAH12 FAH12 WT fad2	0.3	0.3	0.1	0.1
Š	FAH12 WT	0.4	0.4	0.2	0.1
	WT	0	0	0	0
Fatty acid		18:1-ОН	18:2-OH	20:1-OH	20:2-OH

In order to confirm that the observed new peaks in the transgenic lines corresponded to derivatives of ricinoleic, lesquerolic, densipolic and auricolic acids, mass spectrometry was used. The 5 fatty acid derivatives were resolved by gas chromatography as described above except that a Hewlett-Packard 5971 series mass selective detector was used in place of the flame ionization detector used in the previous experiment. The spectra of the four new peaks in Figure 3B (peak numbers 10, 11, 12 10 and 13) are shown in Figures 4A-D, respectively. Comparison of the spectrum obtained for the standards with that obtained for the four peaks from the transgenic lines confirms the identity of the four new peaks. On the basis of the three 15 characteristic peaks at M/Z 187, 270 and 299, peak 10 is unambiguously identified as O-TMSmethylricinoleate. On the basis of the three characteristic peaks at M/Z 185, 270 and 299, peak 11 is unambiguously identified as O-TMS-20 methyldensipoleate. On the basis of the three characteristic peaks at M/Z 187, 298 and 327, peak 12 is unambiguously identified as O-TMSmethyllesqueroleate. On the basis of the three characteristic peaks at M/Z 185, 298 and 327, peak 25 13 is unambiguously identified as O-TMSmethylauricoleate.

These results unequivocally demonstrate the identity of the fahl2 cDNA as encoding a hydroxylase that hydroxylates both oleic acid to produce ricinoleic acid and also hydroxylates icosenoic acid to produce lesquerolic acid. These results also provide additional evidence that the hydroxylase can be functionally expressed in a heterologous plant

species in such a way that the enzyme is catalytically functional. These results also demonstrate that expression of this hydroxylase gene leads to accumulation of ricinoleic, lesquerolic, densipolic and auricolic acids in a plant species that does not normally accumulate hydroxylated fatty acids in extractable lipids.

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The present inventors expected to find lesquerolic acid in the transgenic plants based on the biochemical evidence suggesting broad substrate specificity of the kappa hydroxylase. By contrast, the accumulation of densipolic and auricolic acids was less predictable. Since Arabidopsis does not normally contain significant quantities of the nonhydroxylated precursors of these fatty acids which could serve as substrates for the hydroxylase, it appears that one or more of the three n-3 fatty acid desaturases known in Arabidopsis (e.g., fad3, fad7, fad8; reviewed in Gibson et al., 1995) are capable of desaturating the hydroxylated compounds at the n-3 position. That is, densipolic acid is produced by the action of an n-3 desaturase on ricinoleic acid. Auricolic acid is produced by the action of an n-3 desaturase on lesquerolic acid. Because it is located in the endoplasmic reticulum, the fad3 desaturase is almost certainly responsible. This can be tested in the future by producing fah12containing transgenic plants of the fad3-deficient mutant of Arabidopsis (similar experiments can be done with fad7 and fad8). It is also formally possible that the enzymes that normally elongate 18:1cisa9 to 20:1cisa11 may elongate 120H-18:1cisa9 to $140H-20:1^{c15\Delta11}$, and $120H-18:2^{c15\Delta9,15}$ to $140H-20:2^{c16\Delta11.17}$.

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The amount of the various fatty acids in seed, leaf and root lipids of the control and transgenic plants is also presented in Table 1. Although the amount of hydroxylated fatty acids produced in this example is less than desired for production of ricinoleate and other hydroxylated fatty acids from plants, numerous improvements may be envisioned that will increase the level of accumulation of hydroxylated fatty acids in plants that express the fahl2 or related hydroxylase genes. Improvements in the level and tissue specificity of expression of the hydroxylase gene are envisioned. Methods to accomplish this by the use of strong, seed-specific promoters such as the B. napus napin promoter will be obvious to one skilled in the art. Additional improvements are envisioned that involve modification of the enzymes which cleave hydroxylated fatty acids from phosphatidylcholine, reduction in the activities of enzymes which degrade hydroxylated fatty acids and replacement of acyltransferases which transfer hydroxylated fatty acids to the sn-1, sn-2 and sn-3 positions of glycerolipids. Although genes for these enzymes have not been described in the scientific literature. their utility in improving the level of production of hydroxylated fatty acids can be readily appreciated based on the results of biochemical investigations of ricinoleate synthesis.

Although Arabidopsis is not an economically
important plant species, it is widely accepted by
plant biologists as a model for higher plants.
Therefore, the inclusion of this example is intended
to demonstrate the general utility of the invention
described here to the modification of oil

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composition in higher plants. One advantage of studying the expression of this novel gene in Arabidopsis is the existence in this system of a large body of knowledge on lipid metabolism, as well as the availability of a collection of mutants which 5 can be used to provide useful information on the biochemistry of fatty acid hydroxylation in plant species. Another advantage is the ease of transposing any of the information obtained on 10 metabolism of ricinoleate in Arabidopsis to closely related species such as the crop plants Brassica napus, Brassica juncea or Crambe abyssinica in order to mass produce ricinoleate, lesqueroleate or other hydroxylated fatty acids for industrial use. The 15 kappa hydroxylase is useful for the production of ricinoleate or lesqueroleate in any plant species that accumulates significant levels of the precursors, oleic acid and icosenoic acid. Of particular interest are genetically modified 20 varieties that accumulate high levels of oleic acid. Such varieties are currently available for sunflower and canola. Production of lesquerolic acid and related hydroxy fatty acids can be achieved in species that accumulate high levels of icosenoic 25 acid or other long chain monoenoic acids. Such plants may in the future be produced by genetic engineering of plants that do not normally make such precursors. Thus, the use of the kappa hydroxylase will be of general utility.

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EXAMPLE 2. ISOLATION OF LESQUERELLA KAPPA HYDROXYLASE GENOMIC CLONE Overview

Regions of nucleotide sequence that were conserved in both the castor kappa hydroxylase and the Arabidopsis fad2 $\Delta 12$ fatty acid desaturase were used to design oligonucleotide primers. These were used with genomic DNA from Lesquerella fendleri to amplify fragments of several homologous genes. These amplified fragments were then used as hybridization probes to identify full length genomic clones from a genomic library of L. fendleri.

Hydroxylated fatty acids are specific to the seed tissue of Lesquerella sp., and are not found to any appreciable extent in vegetative tissues. One of the two genes identified by this method was expressed in both leaves and developing seeds and is therefore thought to correspond to the \$\Delta{12}\$ fatty acid desaturase. The other gene was expressed at high levels in developing seeds but was not expressed or was expressed at very low levels in leaves and is the kappa hydroxylase from this species. The identity of the gene as a fatty acyl hydroxylase was established by functional expression of the gene in yeast.

The identity of this gene will also be established by introducing the gene into transgenic Arabidopsis plants and showing that it causes the accumulation of ricinoleic acid, lesquerolic acid, densipolic acid and auricolic acid in seed lipids.

The various steps involved in this process are described in detail below. Unless otherwise indicated, routine methods for manipulating nucleic

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acids, bacteria and phage were as described by Sambrook et al. (1989).

Isolation of a fragment of the Lesquerella kappa hydroxylase gene

Oligonucleotide primers for the amplification of the *L. fendleri* kappa hydroxylase were designed by choosing regions of high deduced amino acid sequence homology between the castor kappa hydroxylase and the *Arabidopsis* Δ12 desaturase (fad2). Because most amino acids are encoded by several different codons, these oligonucleotides were designed to encode all possible codons that could encode the corresponding amino acids.

The sequence of these mixed oligonucleotides was Oligo 1: TAYWSNCAYMGNMGNCAYCA (SEQ ID NO:14) and Oligo 2: RTGRTGNGCNACRTGNGTRTC (SEQ ID NO:15) where Y = C+T, W = A+T, S = G+C, N = A+G+C+T, M = A+C, and R = A+G.

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These oligonucleotides were used to amplify a 20 fragment of DNA from L. fendleri genomic DNA by the polymerase chain reaction (PCR) using the following conditions: Approximately 100 ng of genomic DNA was added to a solution containing 25 pmol of each primer, 1.5 U Taq polymerase (Boehringer Manheim), 200 uM of dNTPs, 50 mM KCl, 10 mM Tris.Cl (pH 9), 25 0.1% (v/v) Triton X-100, 1.5 mM MgCl₂, 3% (v/v)formamide, to a final volume of 50 μ l. Amplifications conditions were: 4 min denaturation step at 94°C, followed by 30 cycles of 92°C for 1 min, 55°C for 1 min, 72°C for 2 min. A final 30 extension step closed the program at 72°C for 5 min.

PCR products of approximately 540 bp were observed following electrophoretic separation of the

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products of the PCR reaction in agarose gels. Two of these fragments were cloned into pBluescript (Stratagene) to give rise to plasmids pLesq2 and pLesq3. The sequence of the inserts in these two plasmids was determined by the chain termination method. The sequence of the insert in pLesq2 is presented as Figure 5 (SEQ ID NO:1) and the sequence of the insert in pLesq3 is presented as Figure 6 (SEQ ID NO:2). The high degree of sequence identity between the two clones indicated that they were both potential candidates to be either a Δ 12 desaturase or a kappa hydroxylase.

Northern analysis

In L. fendleri, hydroxylated fatty acids are found in large amounts in seed oils but are not found in appreciable amounts in leaves. An important criterion in discriminating between a fatty acyl desaturase and kappa hydroxylase is that the kappa hydroxylase gene is expected to be expressed more highly in tissues which have high level of hydroxylated fatty acids than in other tissues. In contrast, all plant tissues should contain mRNA for an ω6 fatty acyl desaturase since diunsaturated fatty acids are found in the lipids of all tissues in most or all plants.

Therefore, it was of great interest to determine whether the gene corresponding to pLesq2 was also expressed only in seeds, or is also expressed in other tissues. This question was addressed by testing for hybridization of pLesq2 to RNA purified from developing seeds and from leaves.

Total RNA was purified from developing seeds and young leaves of L. fendleri using an Rneasy RNA

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extraction kit (Qiagen), according to the manufacturer's instructions. RNA concentrations were quantified by UV spectrophotometry at λ =260 and 280 nm. In order to ensure even loading of the gel to be used for Northern blotting, RNA concentrations were further adjusted after recording fluorescence under UV light of RNA samples stained with ethidium bromide and run on a test denaturing gel.

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Total RNA prepared as described above from leaves and developing seeds was electrophoresed through an agarose gel containing formaldehyde (Iba et al., 1993). An equal quantity (10 μ g) of RNA was loaded in both lanes, and RNA standards (0.16-1.77 kb ladder, Gibco-BRL) were loaded in a third lane. Following electrophoresis, RNA was transferred from the gel to a nylon membrane (Hybond N+, Amersham) and fixed to the filter by exposure to UV light.

A 32 P-labelled probe was prepared from insert DNA of clone pLesq2 by random priming and hybridized to the membrane overnight at 52°C, after it had been prehybridized for 2 h. The prehybridization solution contained 5X SSC, 10X Denhardt's solution, 0.1% SDS, 0.1M KPO₄ pH 6.8, 100 μ g/ml salmon sperm DNA. The hybridization solution had the same basic composition, but no SDS, and it contained 10% dextran sulfate and 30% formamide. The blot was washed once in 2X SSC, 0.5% SDS at 65°C then in 1X SSC at the same temperature.

Brief (30 min) exposure of the blot to X-ray film revealed that the probe pLesq2 hybridized to a single band only in the seed RNA lane (Figure 7). The blot was re-probed with the insert from pLesq3 gene, which gave bands of similar intensity in the seed and leaf lanes (Figure 7).

These results show that the gene corresponding to the clone pLesq2 is highly and specifically expressed in seed of *L. fendleri*. In conjunction with knowledge of the nucleotide and deduced amino acid sequence, strong seed-specific expression of the gene corresponding to the insert in pLesq2 is a convincing indicator of the role of the enzyme in synthesis of hydroxylated fatty acids in the seed oil.

10 <u>Characterization of a genomic clone of the kappa</u> hydroxylase

Genomic DNA was prepared from young leaves of L. fendleri as described by Murray and Thompson (1980). A Sau3AI-partial digest genomic library constructed in the vector \DashII (Stratagene, 11011 15 North Torrey Pines Road, La Jolla CA 92037) was prepared by partially digesting 500 µg of DNA, sizeselecting the DNA on a sucrose gradient (Sambrook et al., 1989), and ligating the DNA (12 kb average 20 size) to the BamHI-digested arms of λDashII. The entire ligation was packaged according to the manufacturer's conditions and plated on E. coli strain XL1-Blue MRA-P2 (Stratagene). This yielded 5x10⁵ primary recombinant clones. The library was then amplified according to the manufacturer's 25 conditions. A fraction of the genomic library was plated on E. coli XL1-Blue and resulting plagues (150,000) were lifted to charged nylon membranes (Hybond N+, Amersham), according to the manufacturer's conditions. DNA was crosslinked to 3 C the filters under UV in a Stratalinker (Stratagene).

Several clones carrying genomic sequences corresponding to the L. fendleri hydroxylase were

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isolated by probing the membranes with the insert from pLesq2 that was PCR-amplified with internal primers and labelled with 32P by random priming. The filters were prehybridized for 2 hours at 65°C in 7% SDS, 1mM EDTA, 0.25 M Na, HPO, (pH 7.2), 1% BSA and hybridized to the probe for 16 hours in the same solution. The filters were sequentially washed at 65°C in solutions containing 2 X SSC, 1 X SSC, 0.5 X SSC in addition to 0.1 % SDS. A 2.6 kb XbaI fragment containing the complete coding sequence for the kappa hydroxylase and approximately 1 kb of the 5' upstream region was subcloned into the corresponding site of pBluescript KS to produce plasmid pLesg-Hyd and the sequence determined completely using an automatic sequencer by the dideoxy chain termination method. Sequence data was analyzed using the program DNASIS (Hitachi Company).

The sequence of the insert in clone pLesq-Hyd is shown in Figures 8A-B. The sequence entails 1855 bp of contiguous DNA sequence (SEQ ID NO:3). The clone encodes a 401 bp 5' untranslated region (i.e., nucleotides preceding the first ATG codon), an 1152 bp open reading frame, and a 302 bp 3' untranslated region. The open reading frame encodes a 384 amino acid protein with a predicted molecular weight of 44,370 (SEQ ID NO:4). The amino terminus lacks features of a typical signal peptide (von Heijne, 1985).

The exact translation-initiation methionine
has not been experimentally determined, but on the
basis of deduced amino acid sequence homology to the
castor kappa hydroxylase (noted below) is thought to
be the methionine encoded by the first ATG codon at
nucleotide 402.

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Comparison of the pLesq-Hyd deduced amino acid sequence with sequences of membrane-bound desaturases and the castor hydroxylase (Figures 9A-B) indicates that pLesq-Hyd is homologous to these genes. This figure shows an alignment of the L. fendleri hydroxylase (SEQ ID NO:4) with the castor hydroxylase (van de Loo et al., 1995), the Arabidopsis fad2 cDNA which encodes an endoplasmic reticulum-localized Δ 12 desaturase (called fad2) (Okuley et al., 1994), two soybean fad2 desaturase clones, a Brassica napus fad2 clone, a Zea mays fad2 clone and partial sequence of a R. communis fad2 clone.

The high degree of sequence homology indicates that the gene products are of similar function. For instance, the overall homology between the Lesquerella hydroxylase and the Arabidopsis fad2 desaturase was 92.2% similarity and 84.8% identity and the two sequences differed in length by only one amino acid.

Southern hybridization

Southern analysis was used to examine the copy number of the genes in the *L. fendleri* genome corresponding to the clone pLesq-Hyd. Genomic DNA (5 μ g) was digested with *Eco*RI, *Hind*III and *Xba*I and separated on a 0.9% agarose gel. DNA was alkaliblotted to a charged nylon membrane (Hybond N+, Amersham), according to the manufacturer's protocol. The blot was prehybridized for 2 hours at 65°C in 7% SDS, 1mM EDTA, 0.25 M Na₂HPO₄ (pH 7.2), 1% BSA and hybridized to the probe for 16 hours in the same solution with pLesq-Hyd insert PCR-amplified with internal primers and labelled with ¹²P by random

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priming. The filters were sequentially washed at 65°C in solutions containing 2 X SSC, 1 X SSC, 0.5 X SSC in addition to 0.1 % SDS, then exposed to X-ray film.

The probe hybridized with a single band in each digest of *L. fendleri* DNA (Figure 10), indicating that the gene from which pLesq-Hyd was transcribed is present in a single copy in the *L. fendleri* genome.

10 Expression of pLesq-Hyd in Transgenic Plants

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There are a wide variety of plant promoter sequences which may be used to cause tissue-specific expression of cloned genes in transgenic plants. For instance, the napin promoter and the acyl carrier protein promoters have previously been used in the modification of seed oil composition by expression of an antisense form of a desaturase (Knutson et al., 1992). Similarly, the promoter for the β subunit of soybean β -conglycinin has been shown to be highly active and to result in tissue-specific expression in transgenic plants of species other than soybean (Bray et al., 1987). Thus, other promoters which lead to seed-specific expression may also be employed for the production of modified seed oil composition. Such modifications of the invention described here will be obvious to one skilled in the art.

Constructs for expression of *L. fendleri* kappa hydroxylase in plant cells are prepared as follows: A 13 kb *SalI* fragment containing the pLesq-Hyg gene was ligated into the *XhoI* site of binary Ti plasmid vector pSLJ44026 (Jones et al., 1992) (Figure 11) to produce plasmid pTi-Hyd and

transformed into Agrobacterium tumefaciens strains GV3101 by electroporation. Strain GV3101 (Koncz and Schell, 1986) contains a disarmed Ti plasmid. Cells for electroporation were prepared as follows. GV3101 was grown in LB medium with reduced NaCl (5 g/l). A 5 250 ml culture was grown to $OD_{600} = 0.6$, then centrifuged at 4000 rpm (Sorvall GS-A rotor) for 15 min. The supernatant was aspirated immediately from the loose pellet, which was gently resuspended in 500 ml ice-cold water. The cells were centrifuged as 10 before, resuspended in 30 ml ice-cold water, transferred to a 30 ml tube and centrifuged at 5000 rpm (Sorvall SS-34 rotor) for 5 min. This was repeated three times, resuspending the cells consecutively in 30 ml ice-cold water, 30 ml ice-15 cold 10% glycerol, and finally in 0.75 ml ice-cold 10% glycerol. These cells were aliquoted, frozen in liquid nitrogen, and stored at -80°C.

Electroporations employed a Biorad Gene

Pulser instrument using cold 2 mm-gap cuvettes containing 40 μl cells and 1 μl of DNA in water, at a voltage of 2.5 KV, and 200 Ohms resistance. The electroporated cells were diluted with 1 ml SOC medium (Sambrook et al., 1989, page A2) and incubated at 28°C for 2-4 h before plating on medium containing kanamycin (50 mg/l).

Arabidopsis thaliana can be transformed with the Agrobacterium cells containing pTi-Hyd as described in Example 1 above. Similarly, the presence of hydroxylated fatty acids in the transgeneic Arabidopsis plants can be demonstrated by the methods described in Example 1 above.

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Constitutive expression of the *L. fendleri* hydroxylase in transgenic plants

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A 1.5 kb EcoRI fragment from pLesg-Hyg comprising the entire coding region of the hydroxylase was gel purified, then cloned into the corresponding site of pBluescript KS (Stratagene). Plasmid DNA from a number of recombinant clones was then restricted with PstI, which should cut only once in the insert and once in the vector polylinker sequence. Release of a 920 bp fragment with PstI indicated the right orientation of the insert for further manipulations. DNA from one such clone was further restricted with SalI, the 5' overhangs filled-in with the Klenow fragment of DNA polymerase I, then cut with SacI. The insert fragment was gel purified, and cloned between the SmaI and SacI sites of pBI121 (Clontech) behind the cauliflower mosaic virus 35S promoter. After checking that the sequence of the junction between insert and vector DNA was appropriate, plasmid DNA from a recombinant clone was used to transform A. tumefaciens (GV3101). Kanamycin resistant colonies were then used for in planta transformation of A. thaliana as previously described.

DNA was extracted from kanamycin resistant seedlings and used to PCR-amplify selected fragments from the hydroxylase using nested primers. When fragments of the expected size could be amplified, corresponding plants were grown in the greenhouse or on agar plates, and fatty acids extracted from fully expanded leaves, roots and dry seeds. GC-MS analysis was then performed as previously described to characterize the different fatty acid species and

detect accumulation of hydroxy fatty acids in transgenic tissues.

Expression of the Lesquerella hydroxylase in yeast

In order to demonstrate that the cloned L. fendleri gene encoded a kappa hydroxylase, the gene was expressed in yeast cells under transcriptional control of an inducible promoter and the yeast cells were examined for the presence of hydroxylated fatty acids by GC-MS.

In a first step, a lambda genomic clone containing the L. fendleri hydroxylase gene was cut with EcoRI, and a resulting 1400 bp fragment containing the coding sequence of the hydroxylase gene was subcloned in the EcoRI site of the pBluescript KS vector (Stratagene). This subclone, pLesqcod, contains the coding region of the Lesquerella hydroxylase plus some additional 3' sequence.

In a second step, pLesqcod was cut with

HindIII and XbaI, and the insert fragment was cloned into the corresponding sites of the yeast expression vector pYes2 (Invitrogen; Figure 12). This subclone, pLesqYes, contains the L. fendleri hydroxylase in the sense orientation relative to the 3' side of the Gall promoter. This promoter is inducible by the addition of galactose to the growth medium, and is repressed upon addition of glucose. In addition, the vector carries origins of replication allowing the propagation of pLesqYes in both yeast and E. coli.

Transformation of S. cerevisiae host strain CGY2557

Yeast strain CGY2557 (MATα, GAL', ura3-52,

leu2-3, trpl, ade2-1, lys2-1, his5, can1-100) was

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grown overnight at 28°C in YPD liquid medium (10 q yeast extract, 20 g bacto-peptone, 20 g dextrose per liter), and an aliquot of the culture was inoculated into 100 ml fresh YPD medium and grown until the OD600 of the culture was 1. Cells were then collected by centrifugation and resuspended in about 200 ul of supernatant. $40\mu l$ aliquots of the cell suspension were then mixed with $1-2\mu q$ DNA and electroporated in 2 mm-gap cuvettes using a Biorad Gene Pulser instrument set at 600 V, 200 Ω , 25 μ F, 160 μ l YPD was added and the cells were plated on selective medium containing glucose. Selective medium consisted of 6.7 q yeast nitrogen base (Difco), 0.4 q casamino acids (Difco), 0.02 g adenine sulfate, 0.03 g Lleucine, 0.02 g L-tryptophan, 0.03 g L-lysine-HCl, 0.03 g L-histidine-HCl , 2% glucose, water to 1 liter. Plates were solidified using 1.5% Difco Bacto-agar. Transformant colonies appeared after 3 to 4 days incubation at 28°C.

Independent transformant colonies from the previous experiment were used to inoculate 5 ml of selective medium containing either 2% glucose (gene repressed) or 2% galactose (gene induced) as the sole carbon source. Independent colonies of CGY2557 transformed with pYES2 containing no insert were used as controls.

After 2 days of growth at 28°C, an aliquot of the cultures was used to inoculate 5 ml of fresh selective medium. The new culture was placed at 16°C and grown for 9 days.

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Fatty acid analysis of yeast expressing the L. fendleri hydroxylase

Cells from 2.5 ml of culture were pelleted at 1800g, and the supernatant was aspirated as completely as possible. Pellets were then dispersed in 1 ml of 1 N methanolic HCl (Supelco, Bellafonte, PA). Transmethylation and derivatization of hydroxy fatty acids were performed as described above. After drying under nitrogen, samples were redissolved in 50µl chloroform before being analyzed by GC-MS. Samples were injected into an SP2330 fused-silica capillary column (30 m x 0.25 mm ID, 0.25µm film thickness, Supelco). The temperature profile was 100 - 160°C, 25°C/min, 160 - 230°C, 10°C/min, 230°C, 3 min, 230-100°C, 25°C/min. Flow rate was 0.9 ml/min. Fatty acids were analyzed using a Hewlett-Packard 5971 series Msdetector.

Gas chromatograms of derivatized fatty acid methyl esters from induced cultures of yeast containing pLesqYes contained a novel peak that eluted at 7.6 min (Figure 13). O-TMS methyl ricinoleate eluted at exactly the same position on control chromatograms. This peak was not present in cultures lacking pLesqYes or in cultures containing pLesqYes grown on glucose (repressing conditions) rather than galactose (inducing conditions). Mass spectrometry of the peak (Figure 13) revealed that the peak has the same spectrum as O-TMS methyl ricinoleate. Thus, on the basis of chromatographic retention time and mass spectrum, it was concluded that the peak corresponded to O-TMS methyl ricinoleate. The presence of ricinoleate in the transgenic yeast cultures confirms the identity of the gene as a kappa hydroxylase of this invention.

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EXAMPLE 3. OBTAINING OTHER PLANT FATTY ACYL HYDROXYLASES

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The castor fahl2 sequence could be used to identify other kappa hydroxylases by methods such as PCR and heterologous hybridization. However, because of the high degree of sequence similarity between $\Delta 12$ desaturases and kappa hydroxylases, the prior art does not teach how to distinguish between the two kinds of enzymes without a functional test such as demonstrating activity in transgenic plants or another suitable host (e.g., transgenic microbial or animal cells). The identification of the L. fendleri hydroxylase provided for the development of criteria by which a hydroxylase and a desaturase may be distinguished solely on the basis of deduced amino acid sequence information.

Figures 9A-B show a sequence alignment of the

castor and L. fendleri hydroxylase sequences with the castor hydroxylase sequence and all publicly available sequences for all plant microsomal $\Delta 12$ 20 fatty acid desaturases. Of the 384 amino acid residues in the castor hydroxylase sequence, more than 95% are identical to the corresponding residue in at least one of the desaturase sequences. 25 Therefore, none of these residues are responsible for the catalytic differences between the hydroxylase and the desaturases. Of the remaining 16 residues in the castor hydroxylase and 14 residues in the Lesquerella hydroxylase, all but seven represent instances where the hydroxylase sequence 30 has a conservative substitution compared with one or more of the desaturase sequences, or there is wide variability in the amino acid at that position in the various desaturases. By conservative, it is

herein.

meant that the following amino acids are functionally equivalent: Ser/Thr, Ile/Leu/Val/Met, Asp/Glu. Thus, these structural differences also cannot account for the catalytic differences between the desaturases and hydroxylases. This leaves just 5 seven amino acid residues where both the castor hydroxylase and the Lesquerella hydroxylase differ from all of the known desaturases and where all of the known microsomal A12 desaturases have the 10 identical amino acid residue. These residues occur at positions 69, 111, 155, 226, 304, 331 and 333 of the alignment in Figure 9. Therefore, these seven sites distinguish hydroxylases from desaturases. Based on this analysis, the present inventors 15 believe that any enzyme with greater than 60% sequence identity to one of the enzymes listed in Figure 9 can be classified as a hydroxylase if it differs from the sequence of the desaturases at these seven positions. Because of slight differences 20 in the number of residues in a particular protein, the numbering may vary from protein to protein but the intent of the number system will be evident if the protein in question is aligned with the castor hydroxylase using the numbering system shown herein. 25 Thus, in conjunction with the methods for using the Lesquerella hydroxylase gene to isolate homologous genes, the structural criterion disclosed here teaches how to isolate and identify plant kappa hydroxylase genes for the purpose of genetically 3 C modifying fatty acid composition as disclosed

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EXAMPLE 4 - USING HYDROXYLASES TO ALTER THE LEVEL OF FATTY ACID UNSATURATION

Evidence that kappa hydroxylases of this invention can be used to alter the level of fatty acid unsaturation was obtained from the analysis of transgenic plants that expressed the castor hydroxylase under control of the cauliflower mosaic virus promoter. The construction of the plasmids and the production of transgenic Arabidopsis plants was described in Example 1 (above). The fatty acid composition of seed lipids from wild type and six transgenic lines (1-2/a, 1-2/b, 1-3/b, 4F, 7E, 7F) is shown in Table 2.

Table 2. Fatty acid composition of lipids from

Arabidopsis seeds. The asterisk (*) indicates that
for some of these samples, the 18:3 and 20:1 peaks
overlapped on the gas chromatograph and, therefore,
the total amount of these two fatty acids is
reported.

TABLE 2	7.F	6	4.2	28.5	19.8	1	2	t	30.6	6.0	9.0	9.0	0.1
	7E	8.4	3.8	30.5	20.1	ı	1.8	,	30.8	0.7	0.5	0.2	0.1
	4 F	8.1	3.5	27.5	21.1	•	2.4	ı	32.1	0.2	0.2	0.2	0
	1-3/b	8.4	3.3	25.5	27.5	14.8	1.1	17.5	ı	0.1	0.1	0	0
	1-2/b	9.5	3.9	34.5	21	14.4	r-t	14.1	ı	0	0	0	0
	1-2/a	9.8	3.8	33	16.9	ı	1.6	1	31.2	9.0	9.0	0.2	0.1
	WT	10.3	3.5	14.7	32.4	13.8	1.3	22.5	1	0	0	0	0
	Fatty acid	16:0	18:0	18:1	18:2	18:3	20:0	20:1	18:3 20:1*	Ricinoleic	Densipolic	Lesquerolic	Auricolic

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The results in Table 2 show that expression of the castor hydroxylase in transgenic Arabidopsis plants caused a substantial increase in the amount of oleic acid (18:1) in the seed lipids and an approximately corresponding decrease in the amount of linoleic acid (18:2). The average amount of oleic acid in the six transgenic lines was 29.9% versus 14.7% in the wild type.

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The precise mechanism by which expression of
the castor hydroxylase gene causes increased
accumulation of oleic acid is not known. However, an
understanding of the mechanism is not required in
order to exploit this invention for the directed
alteration of plant lipid fatty acid composition.

Furthermore, it will be recognized by one skilled in
the art that many improvements of this invention may
be envisioned. Of particular interest will be the
use of other promoters which have high levels of
seed-specific expression.

Since hydroxylated fatty acids were not detected in the seed lipids of transgenic line 1-2b, it seems likely that it is not the presence of hydroxylated fatty acids per se that causes the effect of the castor hydroxylase gene on desaturase activity. Protein-protein interaction between the hydroxylase and the $\Delta 12$ -oleate desaturase or another protein may be required for the overall reaction (e.g., cytochrome b5) or for the regulation of desaturase activity. For example, interaction between the hydroxylase and this other protein may suppress the activity of the desaturase. In particular, the quaternary structure of the membrane-bound desaturases has not been established. It is possible that these enzymes are active as

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dimers or as multimeric complexes containing more than two subunits. Thus, if dimers or multimers form between the desaturase and the hydroxylase, the presence of the hydroxylase in the complex may disrupt the activity of the desaturase.

Transgenic plants may be produced in which the hydroxylase enzyme has been rendered inactive by the elimination of one or more of the histidine residues that have been proposed to bind iron molecules required for catalysis. Several of these histidine residues have been shown to be essential for desaturase activity by site directed mutagenesis (Shanklin et al., 1994). Codons encoding histidine residues in the castor hydroxylase gene will be changed to alanine residues as described by Shanklin et al. (1994). The modified genes will be introduced into transgenic plants of Arabidopsis, and possibly other species such as tobacco, by the methods described in Example 1 of this application.

In order to examine the effect on all tissues, the strong constitutive cauliflower mosaic virus promoter may be used to cause transcription of the modified genes. However, it will be recognized that in order to specifically examine the effect of expression of the mutant gene on seed lipids, a seed-specific promoter such as the B. napus napin promoter may be used. An expected outcome is that expression of the inactive hydroxylase protein in transgenic plants will inhibit the activity of the endoplasmic reticulum-localized $\Delta 12$ -desaturase. Maximum inhibition will be obtained by expressing high levels of the mutant protein.

In a further embodiment of this invention, mutations that inactivate other hydroxylases, such

as the Lesquerella hydroxylase of this invention, may also be useful for decreasing the amount of endoplasmic reticulum-localized $\Delta 12$ -desaturase activity in the same way as the castor gene. In a further embodiment of this invention, similar mutations of desaturase genes may also be used to inactivate endogenous desaturases. Thus, expression of catalytically inactive fad2 gene from Arabidopsis in transgenic Arabidopsis may inhibit the activity of the endogenous fad2 gene product.

Similarly, expression of the catalytically inactive forms of $\Delta 12$ -desaturase from Arabidopsis or other plants in transgenic soybean, rapeseed, Crambe, Brassica juncea, canola, flax, sunflower, safflower, cotton, cuphea, soybean, peanut, coconut, oil palm or corn may lead to inactivation of endogenous $\Delta 12$ -desaturase activity in these plants. In a further embodiment of this invention, expression of catalytically inactive forms of other desaturases such as the $\Delta 15$ -desaturases may lead to inactivation of the corresponding desaturases.

An example of a class of mutants useful in the present invention are "dominant negative" mutants that block the function of a gene at the protein level (Herskowitz, 1987). A cloned gene is altered so that it encodes a mutant product capable of inhibiting the wild type gene product in a cell, thus causing the cell to be deficient in the function of that gene product. Inhibitory variants of a wild type product can be designed because proteins have multiple functional domains that can be mutated independently, e.g., cligomerization, substrate binding, caralysis, membrane association domains or the like. In general, dominant negative

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proteins retain an intact, functional subset of the domains of the parent, wild type protein, but have the complement of that subset either missing or altered so as to be nonfunctional.

Whatever the precise basis for the inhibitory effect of the castor hydroxylase on desaturation, because the castor hydroxylase has very low nucleotide sequence homology (i.e., about 67%) to the Arabidopsis fad2 gene (encoding the endoplasmic reticulum-localized $\Delta 12$ -desaturase), the inhibitory effect of this gene, which is provisionally called "protein-mediated inhibition" ("protibition"), may have broad utility because it does not depend on a high degree of nucleotide sequence homology between the transgene and the endogenous target gene. In particular, the castor hydroxylase may be used to inhibit the endoplasmic reticulum-localized $\Delta 12$ desaturase activity of all higher plants. Of particular relevance are those species used for oil production. These include but are not limited to rapeseed, Crambe, Brassica juncea, canola, flax, sunflower, safflower, cotton, cuphea, soybean, peanut, coconut, oil palm and corn.

CONCLUDING REMARKS

By the above examples, demonstration of critical factors in the production of novel hydroxylated fatty acids by expression of a kappa hydroxylase gene from castor in transgenic plants is described. In addition, a complete cDNA sequence of the Lesquerella fendleri kappa hydroxylase is also provided. A full sequence of the castor hydroxylase is also given with various constructs for use in host cells. Through this invention, one can obtain

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the amino acid and nucleic acid sequences which encode plant fatty acyl hydroxylases from a variety of sources and for a variety of applications. Also revealed is a novel method by which the level of fatty acid desaturation can be altered in a directed way through the use of genetically altered hydroxylase or desaturase genes.

All publications mentioned in this specification are indicative of the level of skill of those skilled in the art to which this invention pertains. All publications are herein incorporated by reference to the same extent as if each individual publication was specifically and individually indicated to be incorporated by reference.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be obvious that certain changes and modifications may be practiced within the scope of the appended claims.

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1	(1)	GENERAL	INFORMATION:
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(i) APPLICANT:

(/	ODIVERGE	THE OWNER FOR .	

Broun, Pierre

van de Loo, Frank

Somerville, Chris

Boddupalli, Sekhar S.

- (ii) TITLE OF INVENTION: Production of Hydroxylated Fatty Acids in Genetically Modified Plants
- (iii) NUMBER OF SEQUENCES: 15
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 - (E) COUNTRY: USA
 - (F) ZIP: 20005-3918
- COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: 3.5 inch, 1.44 MB storage

 - (B) COMPUTER: IBM compatible (C) OPERATING SYSTEM: DOS 5.0
 - (D) SOFTWARE: Word Perfect 5.1
- (vi) CURRENT APPLICATION DATA;
 - (A) APPLICATION NUMBER: not yet assigned
 - FILING DATE: February 6, 1997 (B)
 - (C) CLASSIFICATION:
- (2) INFORMATION FOR SEQ ID NO:1
- SEQUENCE CHARACTERISTICS: (i)
 - (A) LENGTH: 543 nucleotides
 - (B) TYPE: nucleotide
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

40	GGATCCCTAG	TTCCAACAAT	GGCGGCACCA	TATTGGCACC
80	AAGCTGCAGT	CCACCTAAGA	AGTCTTTGTC	AAAAAGATGA
120	TCTTGGACGC	TCAACAACCC	GTCAAATACC	CANATGGTAT
160	GGGTGGCCTT	GTTTATCCTC	TAACAGTTCA	ATTCTGGTGT

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TGTATCTAGC CTTTAATGTA TCAGGTAGAC CTTATGATGG	200
TTTCGCTTCA CATTTCTTCC CTCATGCACC TATCTTTAAG	240
GACCGTGAAC GTCTCCAGAT ATACATCTCA GATGCTGGTA	280
TTCTAGCTGT CTGTTATGGT CTTTACCGTT ACGCTGCTTC	320
ACAAGGATTG ACTGCTATGA TCTGCGTCTA CGGAGTACCG	360
CTTTTGATAG TGAACTTTTT CCTTGTCTTG GTCACTTTCT	400
TGCAGCACAC TCATCCTTCA TTACCTCACT ATGATTCAAC	440
CGAGTGGGAA TGGATTAGAG GAGCTTTGGT TACGGTAGAC	480
AGAGACTATG GAATCTTGAA CAAGGTGTTT CACAACATAA	520
CAGACACCCA CGTAGCACAC CAC	543
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 544 nucleotides (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2: 	
TATAGGCACC GGAGGCACCA TTCCAACACA GGATCCCTCG	40
AAAGAGATGA AGTATTTGTC CCAAAGCAGA AATCCGCAAT	80
CAAGTGGTAC GGCGAATACC TCAACAACCC TCCTGGTCGC	120
ATCATGATGT TAACTGTCCA GTTCGTCCTC GGATGGCCCT	160
TGTACTTAGC CTTCAACGTT TCTGGCAGAC CCTACAATGG	200
TTTCGCTTCC CATTTCTTCC CCAATGCTCC TATCTACAAC	240
GACCGTGAAC GCCTCCAGAT TTACATCTCT GATGCTGGTA	280
TTCTAGCCGT CTGTTATGGT CTTTACCGTT ACGCTGTTGC	320
ACAAGGACTA GCCTCAATGA TCTGTCTAAA CGGAGTTCCG	360
CTTCTGATAG TTAACTTTTT CCTCGTCTTG ATCACTTACT	400

TACAACACA TCACCCTGCG TTGCCTCACT ATGATTCATC	440
AGAGTGGGAT TGGCTTAGAG GAGCTTTAGC TACTGTAGAC	480
AGAGACTATG GAATCTTGAA CAAGGTGTTC CATAACATCA	520
CAGACACCCA CGTCGCACAC CACT	544
(2) INFORMATION FOR SEQ ID NO:3	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1855 nucleotides (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
ATGAAGCTTT ATAAGAAGTT AGTTTTCTCT GGTGACAGAG	40
AAATTNTGTC AATTGGTAGT GACAGTTGAA GCAACAGGAA	80
CAACAAGGAT GGTTGGTGNT GATGCTGATG TGGTGATGTG	120
TTATTCATCA AATACTAAAT ACTACATTAC TTGTTGCTGC	160
CTACTTCTCC TATTTCCTCC GCCACCCATT TTGGACCCAC	200
GANCCTTCCA TTTAAACCCT CTCTCGTGCT ATTCACCAGA	240
AGAGAAGCCA AGAGAGAGA AGAGAGAATG TTCTGAGGAT	280
CATTGTCTTC TTCATCGTTA TTAACGTAAG TTTTTTTGA	320
CCACTCATAT CTAAAATCTA GTACATGCAA TAGATTAATG	360
ACTGTTCCTT CTTTTGATAT TTTCAGCTTC TTGAATTCAA	400
GATGGGTGCT GGTGGAAGAA TAATGGTTAC CCCCTCTTCC	440
AAGAAATCAG AAACTGAAGC CCTAAAACGT GGACCATGTG	480
AGAAACCACC ATTCACTGTT AAAGATCTGA AGAAAGCAAT	520
CCCACAGCAT TGTTTCAAGC GCTCTATCCC TCGTTCTTTC	560
TCCTACCTTC TCACAGATAT CACTTTAGTT TCTTGCTTCT	600

ACTACGTTGC CACAAATTAC TTCTCTCTTC TTCCTCAGCC

TCTCTCTACT	TACCTAGCTT	GGCCTCTCTA	TTGGGTATGT	680
CAAGGCTGTG	TCTTAACCGG	TATCTGGGTC	ATTGGCCATG	720
AATGTGGTCA	CCATGCATTC	AGTGACTATC	AATGGGTAGA	760
TGACACTGTT	GGTTTTATCT	TCCATTCCTT	CCTTCTCGTC	800
CCTTACTTCT	CCTGGAAATA	CAGTCATCGT	CGTCACCATT	840
CCAACAATGG	ATCTCTCGAG	AAAGATGAAG	TCTTTGTCCC	880
ACCGAAGAAA	GCTGCAGTCA	AATGGTATGT	TAAATACCTC	920
AACAACCCTC	TTGGACGCAT	TCTGGTGTTA	ACAGTTCAGT	960
TTATCCTCGG	GTGGCCTTTG	TATCTAGCCT	TTAATGTATC	1000
AGGTAGACCT	TATGATGGTT	TCGCTTCACA	TTTCTTCCCT	1040
CATGCACCTA	TCTTTAAAGA	CCGAGAACGC	CTCCAGATAT	1080
ACATCTCAGA	TGCTGGTATT	CTAGCTGTCT	GTTATGGTCT	1120
TTACCGTTAC	GCTGCTTCAC	AAGGATTGAC	TGCTATGATC	1160
TGCGTCTATG	GAGTACCGCT	TTTGATAGTG	AACTTTTTCC	1200
TTGTCTTGGT	AACTTTCTTG	CAGCACACTC	ATCCTTCGTT	1240
ACCTCATTAT	GATTCAACCG	AGTGGGAATG	GATTAGAGGA	1280
GCTTTGGTTA	CGGTAGACAG	AGACTATGGA	ATATTGAACA	1320
AGGTGTTCCA	TAACATAACA	GACACACATG	TGGCTCATCA	1360
TCTCTTTGCA	ACTATACCGC	ATTATAACGC	AATGGAAGCT	1400
ACAGAGGCGA	TAAAGCCAAT	ACTTGGTGAT	TACTACCACT	1440
TCGATGGAAC	ACCGTGGTAT	GTGGCCATGT	ATAGGGAAGC	1480
AAAGGAGTGT	CTCTATGTAG	AACCGGATAC	GGAACGTGGG	1520
AAGAAAGGTG	TCTACTATTA	CAACAATAAG	TTATGAGGCT	1560
GATAGGGCGA	GAGAAGTGCA	ATTATCAATC	TTCATTTCCA	1600
TGTTTTAGGT	GTCTTGTTTA	AGAAGCTATG	CTTTGTTTCA	1640
ATAATCTCAG	AGTCCATNTA	CTTGTGTTCT	GGTGCATTTT	1680

GCC	TAGT	TAT	GTGG	TGTC	GG A	AGTT	AGTG	т тс	AAACTGCT		
TCC'	TGCT	GTG ·	CTGC	CCAG	TG A	AGAA	CAAG	T TT	ACGTGTTT		
AAA	ATAC'	TCG (GAAC	GAAT	TG A	CCAC	AANA	T AT	CCAAAACC		
GGC'	GGCTATCCGA ATTCCATATC CGAAAACCGG ATATCCAAAT										
TTC	TTCCAGAGTA CTTAG										
(2)	I	VFORI	ITAN	ON F	OR SI	EQ I	D NO	: 4			
(i)	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 384 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (D) TOPOLOGY: linear										
(xi)	SEC	QUENC	CE DI	ESCR	IPTIC	ON:	SEQ	ID I	NO:4:		
Met	Gly	Ala	Gly	Gly 5	Arg	Ile	Met	Val	Thr 10		
Pro	Ser	Ser	Lys	Lys 15	Ser	Glu	Thr	Glu	Ala 20		
Leu	Lys	Arg	Gly	Pro 25	Cys	Glu	Lys	Pro	Pro 30		
Phe	Thr	Val	Lys	Asp 35	Leu	Lys	Lys	Ala	Ile 40		
Pro	Gln	His	Cys	Phe 45	Lys	Arg	Ser	Ile	Pro 50		
Arg	Ser	Phe	Ser	Tyr 55	Leu	Leu	Thr	Asp	Ile 60		
Thr	Leu	Val	Ser	Cys 65	Phe	Tyr	Tyr	Val	Ala 70		
Thr	Asn	Tyr	Phe	Ser 75	Leu	Leu	Pro	Gln	Pro 80		
Leu	Ser	Thr	Tyr	Leu 85	Ala	Trp	Pro	Leu	Tyr 90		
Trp	Val	Cys	Gln	Gly 95	Cys	Val	Leu	Thr	Gly 100		

Ile	Trp	Val	Ile	Gly 105	His	Glu	Cys	Gly	His 110
His	Ala	Phe	Ser	Asp 115	Tyr	Gln	Trp	Val	Asp 120
Asp	Thr	Val	Gly	Phe 125	Ile	Phe	His	Ser	Phe 130
Leu	Leu	Val	Pro	Tyr 135	Phe	Ser	Trp	Lys	Tyr 140
Ser	His	Arg	Arg	His 145	His	Ser	Asn	Asn	Gly 150
Ser	Leu	Glu	Lys	Asp 155	Glu	Val	Phe	Val	Pro 160
Pro	Lys	Lys	Ala	Ala 165	Val	Lys	Trp	Tyr	Val 170
Lys	Tyr	Leu	Asn	As n 175	Pro	Leu	Gly	Arg	Ile 180
Leu	Val	Leu	Thr	Val 185	Gln	Phe	Ile	Leu	Gly 190
Trp	Pro	Leu	Tyr	Leu 195	Ala	Phe	Asn	Val	Ser 200
Gly	Arg	Pro	Tyr	Asp 205	Gly	Phe	Ala	Ser	His 210
Phe	Phe	Pro	His	Ala 215	Pro	Ile	Phe	Lys	Asp 220
Arg	Glu	Arg	Leu	Gln 225	Ile	Tyr	Ile	Ser	Asp 230
Ala	Gly	Ile	Leu	Ala 235	Val	Cys	Tyr	Gly	Leu 240
Tyr	Arg	Tyr	Ala	Ala 245	Ser	Gln	Gly	Leu	Thr 250
Ala	. Met	Ile	Cys	Val 255	Tyr	Gly	Val	Pro	Leu 260
Leu	lle	Val	Asn	Phe 265	Phe	Leu	Val	Leu	Val 270

Thr	Phe	Leu	Gln	His 275	Thr	His	Pro	Ser	Leu 280
Pro	His	Tyr	Asp	Ser 285	Thr	Glu	Trp	Glu	Trp 290
Ile	Arg	Gly	Ala	Leu 295	Val	Thr	Val	Asp	Arg 300
Asp	Tyr	Gly	Ile	Leu 305	Asn	Lys	Val	Phe	His 310
Asn	Ile	Thr	Asp	Thr 315	His	Val	Ala	His	His 320
Leu	Phe	Ala	Thr	Ile 325	Pro	His	Tyr	Asn	Ala 330
Met	Glu	Ala	Thr	Glu 335	Ala	Ile	Lys	Pro	Ile 340
Leu	Gly	Asp	Tyr	Tyr 345	His	Phe	Asp	Gly	Thr 350
Pro	Trp	Tyr	Val	Ala 355	Met	Tyr	Arg	Glu	Ala 360
Lys	Glu	Cys	Leu	Tyr 365	Val	Glu	Pro	Asp	Thr 370
Glu	Arg	Gly	Lys	Lys 375	Gly	Val	Tyr	Tyr	Tyr 380
Asn	Asn	Lys	Leu						

- (2) INFORMATION FOR SEQ ID NO:5
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 387 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Gly Gly Gly Arg Met Ser Thr Val

Ile Thr Ser Asn Asn Ser Glu Lys Lys Gly
15 20

Gly	Ser	Ser	His	Leu 25	Lys	Arg	Ala	Pro	His 30
Thr	Lys	Pro	Pro	Phe 35	Thr	Leu	Gly	Asp	Leu 40
Lys	Arg	Ala	Ile	Pro 45	Pro	His	Cys	Phe	Glu 50
Arg	Ser	Phe	Val	Arg 55	Ser	Phe	Ser	Tyr	Val 60
Ala	Tyr	Asp	Val	Cys 65	Leu	Ser	Phe	Leu	Phe 70
Tyr	Ser	Ile	Ala	Thr 75	Asn	Phe	Phe	Pro	Tyr 80
Ile	Ser	Ser	Pro	Leu 85	Ser	Tyr	Val	Ala	Trp 90
Leu	Val	Tyr	Trp	Leu 95	Phe	Gln	Gly	Cys	Ile 100
Leu	Thr	Gly	Leu	Trp 105	Val	Ile	Gly	His	Glu 110
Cys	Gly	His	His	Ala 115	Phe	Ser	Glu	Tyr	Gln 120
Leu	Ala	Asp	Asp	Ile 125	Val	Gly	Leu	Ile	Val 130
His	Ser	Ala	Leu	Leu 135	Val	Pro	Tyr	Phe	Ser 140
Trp	Lys	Tyr	Ser	His 145	Arg	Arg	His	His	Ser 150
Asn	Ile	Gly	Ser	Leu 155	Glu	Arg	Asp	Glu	Val 160
Phe	Val	Pro	Lys	Ser 165	Lys	Ser	Lys	Ile	Ser 170
Trp	Tyr	Ser	Lys	Tyr 175	Ser	Asn	Asn	Pro	Pro 180
Gly	Arg	Val	Leu	Thr 185	Leu	Ala	Ala	Thr	Leu 190

Leu	Leu	Gly	Trp	Pro 195	Leu	Tyr	Leu	Ala	Phe 200
Asn	Val	Ser	Gly	Arg 205	Pro	Tyr	Asp	Arg	Phe 210
Ala	Cys	His	Tyr	Asp 215	Pro	Tyr	Gly	Pro	Ile 220
Phe	Ser	Glu	Arg	Glu 225	Arg	Leu	Gln	Ile	Tyr 230
Ile	Ala	Asp	Leu	Gly 235	Ile	Phe	Ala	Thr	Thr 240
Phe	Val	Leu	Tyr	Gln 245	Ala	Thr	Met	Ala	Lys 250
Gly	Leu	Ala	Trp	Val 255	Met	Arg	Ile	Tyr	Gly 260
Val	Pro	Leu	Leu	Ile 265	Val	Asn	Cys	Phe	Leu 270
Val	Met	Ile	Thr	Tyr 275	Leu	Gln	His	Thr	His 280
Pro	Ala	Ile	Pro	Arg 285	Tyr	Gly	Ser	Ser	Glu 290
Trp	Asp	Trp	Leu	Arg 295	Gly	Ala	Met	Val	Thr 300
Val	Asp	Arg	Asp	Tyr 305	Gly	Val	Leu	Asn	Lys 310
Val	Phe	His	Asn	Ile 315	Ala	Asp	Thr	His	Val 320
Ala	His	His	Leu	Phe 325	Ala	Thr	Val	Pro	His 330
Tyr	His	Ala	Met	Glu 335	Ala	Thr	Lys		Ile 340
Lys	Pro	Ile	Met	Gly 345	Glu	Tyr	Tyr	Arg	Tyr 350
Asp	Gly	Thr	Pro	Phe 355	Tyr	Lys	Ala	Leu	Trp 360

Arg Glu Ala Lys Glu Cys Leu Phe Val Glu 365 370

Pro Asp Glu Gly Ala Pro Thr Gln Gly Val 375 380

- Phe Trp Tyr Arg Asn Lys Tyr 385
- (2) INFORMATION FOR SEQ ID NO:6
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 383 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Met Gly Ala Gly Gly Arg Met Pro Val Pro
5 10

Thr Ser Ser Lys Lys Ser Glu Thr Asp Thr
15 20

Thr Lys Arg Val Pro Cys Glu Lys Pro Pro 25

Phe Ser Val Gly Asp Leu Lys Lys Ala Ile 35 40

Pro Pro His Cys Phe Lys Arg Ser Ile Pro 45 50

Arg Ser Phe Ser Tyr Leu Ile Ser Asp Ile
55 60

Ile Ile Ala Ser Cys Phe Tyr Tyr Val Ala 65 70

Thr Asn Tyr Phe Ser Leu Leu Pro Gln Pro
75 80

Leu Ser Tyr Leu Ala Trp Pro Leu Tyr Trp 85 90

Ala Cys Gln Gly Cys Val Leu Thr Gly Ile 95 100

Trp Val Ile Ala His Glu Cys Gly His His 105 110

Ala	Phe	Ser	Asp	Tyr 115	Gln	Trp	Leu	Asp	Asp 120
Thr	Val	Gly	Leu	Ile 125	Phe	His	Ser	Phe	Leu 130
Leu	Val	Pro	Tyr	Phe 135	Ser	Trp	Lys	Tyr	Ser 140
His	Arg	Arg	His	His 145	Ser	Asn	Thr	Gly	Ser 150
Leu	Glu	Arg	Asp	Glu 155	Val	Phe	Val	Pro	Lys 160
Gln	Lys	Ser	Ala	Ile 165	Lys	Trp	Tyr	Gly	Lys 170
Tyr	Leu	Asn	Asn	Pro 175	Leu	Gly	Arg	Ile	Met 180
Met	Leu	Thr	Val	Gln 185	Phe	Val	Leu	Gly	Trp 190
Pro	Leu	Tyr	Leu	Ala 195	Phe	Asn	Val	Ser	Gly 200
Arg	Pro	Tyr	Asp	Gly 205	Phe	Ala	Cys	His	Phe 210
Phe	Pro	Asn	Ala	Pro 215	Ile	Tyr	Asn	Asp	Arg 220
Glu	Arg	Leu	Gln	Ile 225	Tyr	Leu	Ser	Asp	Ala 230
Gly	Ile	Leu	Ala	Val 235	Cys	Phe	Gly	Leu	Tyr 240
Arg	Tyr	Ala	Ala	Ala 245	Gln	Gly	Met	Ala	Ser 250
Met	Ile	Cys	Leu	Tyr 255	Gly	Val	Pro	Leu	Leu 260
Ile	Val	Asn	Ala	Phe 265	Leu	Val	Leu	Ile	Thr 270
Tyr	Leu	Gln	His	Thr 275	His	Pro	Ser	Leu	Pro 280

His	Tyr	Asp	Ser	Ser 285	Glu	Trp	Asp	Trp	Leu 290
Arg	Gly	Ala	Leu	Ala 295	Thr	Val	Asp	Arg	Asp 300
Tyr	Gly	Ile	Leu	Asn 305	Lys	Val	Phe	His	Asn 310
Ile	Thr	Asp	Thr	His 315	Val	Ala	His	His	Leu 320
Phe	Ser	Thr	Met	Pro 325	His	Tyr	Asn	Ala	Met 330
Glu	Ala	Thr	Lys	Ala 335	Ile	Lys	Pro	Ile	Leu 340
Gly	Asp	Tyr	Tyr	Gln 345	Phe	Asp	Gly	Thr	Pro 350
Trp	Tyr	Val	Ala	Met 355	Tyr	Arg	Glu	Ala	Lys 360
Glu	Cys	Ile	Tyr	Val 365	Glu	Pro	Asp	Arg	Glu 370
Gly	Asp	Lys	Lys	Gly 375	Val	Tyr	Trp	Tyr	As n 380
Asn	Lys	Leu							

- (2) INFORMATION FOR SEQ ID NO:7
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 384 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:
- Met Gly Ala Gly Gly Arg Met Gln Val Ser 5
- Pro Pro Ser Lys Lys Ser Glu Thr Asp Asn 15 20
- Ile Lys Arg Val Pro Cys Glu Thr Pro Pro 25 30

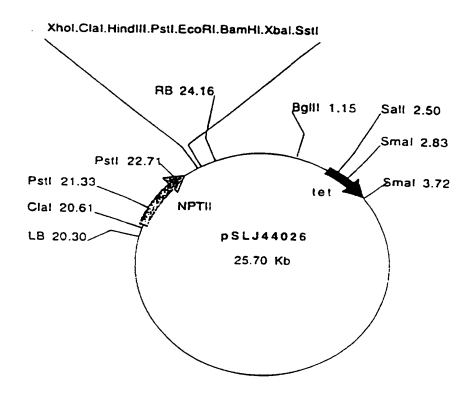
Phe	Thr	Val	Gly	Glu 35	Leu	Lys	Lys	Ala	Ile 40
Pro	Pro	His	Cys	Phe 45	Lys	Arg	Ser	Ile	Pro 50
Arg	Ser	Phe	Ser	His 55	Leu	Ile	Trp	Asp	Ile 60
Ile	Ile	Ala	Ser	Cys 65	Phe	Tyr	Tyr	Val	Ala 70
Thr	Thr	Tyr	Phe	Pro 75	Leu	Leu	Pro	Asn	Pro 80
Leu	Ser	Tyr	Phe	Ala 85	Trp	Pro	Leu	Tyr	Trp 90
Ala	Cys	Gln	Gly	Cys 95	Val	Leu	Thr	Gly	Val 100
Trp	Val	Ile	Ala	His 105	Glu	Cys	Gly	His	Ala 110
Ala	Phe	Ser	Asp	Tyr 115	Gln	Trp	Leu	Asp	Asp 120
Thr	Val	Gly	Leu	Ile 125	Phe	His	Ser	Phe	Leu 130
Leu	Val	Pro	Tyr	Phe 135	Ser	Trp	Lys	Tyr	Ser 140
His	Arg	Arg	His	His 145	Ser	Asn	Thr	Gly	Ser 150
Leu	Glu	Arg	Asp	Glu 155	Val	Phe	Val	Pro	Arg 160
Arg	Ser	Gln	Thr	Ser 165	Ser	Gly	Thr	Ala	Ser 170
Thr	Ser	Thr	Thr	Phe 175	Gly	Arg	Thr	Val	Me t 180
Leu	Thr	Val	Gln	Phe 185	Thr	Leu	Gly	Trp	Pro 190
Leu	Tyr	Leu	Ala	Phe 195	Asn	Val	Ser	Gly	Arg 200

Pro	Tyr	Asp	Gly	Gly 205	Phe	Ala	Cys	His	Phe 210
His	Pro	Asn	Ala	Pro 215	Ile	Tyr	Asn	Asp	Arg 220
Glu	Arg	Leu	Gln	Ile 225	Tyr	Ile	Ser	Asp	Ala 230
Gly	Ile	Leu	Ala	Val 235	Cys	Tyr	Gly	Leu	Leu 240
Pro	Tyr	Ala	Ala	Val 245	Gln	Gly	Val	Ala	Ser 250
Met	Val	Cys	Phe	Leu 255	Arg	Val	Pro	Leu	Leu 260
Ile	Val	Asn	Gly	Phe 265	Leu	Val	Leu	Ile	Thr 270
Tyr	Leu	Gln	His	Thr 275	His	Pro	Ser	Leu	Pro 280
His	Tyr	Asp	Ser	Ser 285	Glu	Trp	Asp	Trp	Leu 290
Arg	Gly	Ala	Leu	Ala 295	Thr	Val	Asp	Arg	Asp 300
Tyr	Gly	Ile	Leu	Asn 305	Gln	Gly	Phe	His	Asn 310
Ile	Thr	Asp	Thr	His 315	Glu	Ala	His	His	Leu 320
Phe	Ser	Thr	Met	Pro 325	His	Tyr	His	Ala	Met 330
Glu	Ala	Thr	Lys	Ala 335	Ile	Lys	Pro	Ile	Leu 340
Gly	Glu	Tyr	Tyr	Gln 345	Phe	Asp	Gly	Thr	Pro 350
Val	Val	Lys	Ala	Met 355	Trp	Arg	Glu	Ala	Lys 360
Glu	Cys	Ile	Tyr	Va l 365	Glu	Pro	Asp	Arg	Gln 370

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/02187

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A01H 5/00, 5/10; C12N 15/52; 15/82 US CL :800/205; 435/172.3, 419; 536/23.6 According to International Patent Classification (IPC) or to both national classification and IPC								
B. FIELDS SEARCHED								
Minimum documentation searched (classification system followe	d by classification symbols)							
U.S. : 800/205; 435/172.3, 419; 536/23.6								
Documentation searched other than minimum documentation to th	e extent that such documents are included in	the fields searched						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) APS, DIALOG								
C. DOCUMENTS CONSIDERED TO BE RELEVANT								
Category* Citation of document, with indication, where a	opropriate, of the relevant passages	Relevant to claim No.						
X WO 94/11516 A1 (LIGHTNER especially pages 40-44 and 109.	1	1, 3, 7, 10, 11, 16, 17, 22, 23, 27-34						
		2, 4-9, 12-15, 18, 21, 24-26						
Further documents are listed in the continuation of Box C	See patent family annex.							
Special categories of cited documents:	"?" later document published after the inserna date and not in conflict with the application	stional filing date or priority						
"A" document defining the general state of the art which is not considered to be of particular relevance	principle or theory underlying the inventi	00						
*E" earlier document published on or after the international filing date	"X" document of particular relevance; the ci considered novel or cannot be considered	laimed invention cannot be to involve an inventive sten						
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	when the document is taken alone "Y" document of particular relevance; the cl	laimed invention cannot be						
O document referring to an oral disclosure, use, exhibition or other means	considered to involve an inventive ste combined with one or more other such do being obvious to a person skilled in the a	ocuments, such combination						
"P" document published prior to the international filing date but later than the priority date claimed	*&* document member of the same patent fam	nil						
Date of the actual completion of the international search	Date of mailing of the international search	h report						
03 JUNE 1997	30 JUL 1997							
Name and mailing address of the ISA/US	Authorized officer							
Commissioner of Patents and Trademarks Box PCT	ELIZABETH F. MCELWAIN	Y						
Washington, D.C. 20231 Facsimile No. (703) 305-3230	Telephone No. (703) 308-0196							



Plasmid name: pSLJ44026 Plasmid size: 25.70 kb

Constructed by: Jonathon Jones

Construction date: 1992

Comments/References: Transgenic Research 1,285-297 (1992)

Figure 11

GA EcoRI HindIII

3.6 -

1.8 -

1.5 -

FIG. 10

			•				
		260	270	280	290	300	
LFFAH12		YAASQGLTAM			LOHTHPSLPH		300
FAH12		ATMAKGLAWV			LOHTHPAIPR		300
ATFAD2	251	YAAAQGMASM	ICLYGVPLLI		LOHTHPSLPH		300
BNFAD2	251	YAAVQGVASM			LOHTHPSLPH		300
GMFAD2-1	251		LCVYGVPLLI		LOHTHFALPH		300
GMFAD2-2		LAMAKGLAWV	VCVYGVPLLV		LOHTHPALPH		300
ZMFAD2	251	LAAAFGVWWV	VRVYAVPLLI		LOHTHPSLPH		300
RCFAD2	251	LAIAKGLAWV	VCVYGVPLLV	VNSFLVLITF	LOHTHPALPH	YDSSEWDWLR	300
15514110		310	320		340	350	
LFFAH12			GILNKVFHNI		ATIPHYNAME		350
FAH12	301	GAMVTVDRDY			ATVPHYHAME		350
ATFAD2	301	GALATVORDY			STMPHYNAME		350
BNFAD2	301	GALATVDRDY	GILNOGFHNI				350
GMFAD2-1	301	GALATMDRDY	GILNKVFHHI		STMPHYHAME		350
GMFAD2 - 2	301	GALATVDRDY			STMPHYHAME		350
ZMFAD2	301	GALATMORDY		TOTHVAHHLF	STMPHYHAME	ATKAIRPILG	350
RCFAD2	301		GILNKVFHNI		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • •	350
1.554411.0	253	360	370	380	390	400	
LFFAH12	351	DYYHFDGTPW	YVAMYREAKE	CLYVEPDTER	GKKGVYYYNN	K-L	400
FAH12	351		YKALWREAKE				400
ATFAD2	351		YVAMYREAKE		DKKGVYWYNN		400
BNFAD2	351		VKAMWREAKE		EKKGVFWYNN		400
GMFAD2-1	351		YKALWREARE			KY	400
GMFAD2-2	351		VKAMWREARE				400
ZMFAD2	351	UYYHFDPTPV	AKATWREAGE	CIYVEPE	DRKGVFWYNK	KF*	400

Figure 9B

		10	. 20	0 3	0 4	0 50	
LFFAH1	2	MGAGGRIM-	VTPSSKKS	SETEALKR	C DEEKDBETY	V 01 444 1 100 10	.
FAH12	-	I CHADOORHS I I	/ 1 2 N N 2 E K K (•GSSHLKR	A PHTKPPFTI	C DINDYIDDAC	50
ATFAD2	•	TIGAGGRIP -	A L 1 2 2 K K 3	SEIDITKR	V PCFKPPFSV	C DIKKYTOBUC	50
BNFAD2		MGAGGRMO-		FINNIKD'	V DEETDDEETW	C ELVULTABLE	50
GMFAD2		MGLA-KETTN	I GUKUKVAKVE	: VUGKKPLSR'	V PNTKPPFTV	C ULKAYIDOUC	50
GMFAD2	·2]	ייי אטטאטויי	TUVPPANKKS	FVDPIKRI	/ DEEKDUECT	C ATPAIRMEN 2	50
ZMFAD2]	MGAGGRMTER	EREKOEOLAR	RATGGAAMOR	DVEVDOET!	3 GIKKAIPPHC	50
RCFAD2	1		• • • • • • • • •	· ··········		3 GIKKAIPPHC	50
		60	70	90) 0		50
LFFAH12	51	FKRSIPRSES	YIITOTTIVS	CEVVVATAVI	9(5 ST BODI 67)) 100 (LAWPLYWVCQ	
FAH12	51	FERSEVESES	YVAYDVCISE	I EVCTATMEN	SELPUPLSI	Y VAWLVYWLFQ	100
ATFAD2	51	FKRSIPRSES	YITOOTTIAS	CITOLATIVE	L L11225F2-1	Y VAWLVYWLFQ	100
BNFAD2	51	FKRSTPRSES	HIJUNITAS	CEVVUATTO	200000000	' FAWPLYWACQ	100
GMFAD2 -		FORSILISES	AAAADIIIVO	TEV. TATTUR	PLLPNPLS-Y	FAWPLYWACO.	100
GMFAD2-		FORSVIRSES	VVVVDCSFAF	TETTLATIVE	HLLPOPFS-L	IAWPIYWVLQ	100
ZMFAD2		FORSVLRSFS	AAAAACITAA	CLITVAINTE	HLLPGPLS	RGMAIYWAVQ	100
RCFAD2	51	FERSVLKSFS	IAAUDEATAY	ALLYFALAII	PALPSPLR-Y	AAWPLYWIAQ	100
	31		120	••••••	• • • • • • • • •	• • • • • • • • •	100
LFFAH12	101	110	120	130	140	150	
FAH12		GCVLTGIWVI	CHECCHHAFS	DYOWVDDIVG	FIFHSFLLVP	YFSWKYSHRR	150
ATFAD2	101	GCILTGLWVI	AMECGHHAFS	EYULADDIVG	LIVHSALLVP	YFSWKYSHRR	150
BNFAD2	101	GCVLTGIWVI	AHECGHHAFS	DYOWLDDIVG	LIFHSFLLVP	YFSWKYSHRR	150
GMFAD2-	1 101	GCVLTGVWVI	AHECGHAAFS	DYOWLDDTVG	LIFHSFLLVP	YFSWKYSHRR	150
GMFAD2-	1 101	GLLLIGVWVI	AHFI GHMAFC	EXULIVADOUS	_ Tiberiio	14 = 0	150
ZMFAD2	c ivi	GCILIGVWVI	AMPLIGHMAF	DAULTUULAG	I TIUCALLUO	VECLUME	150
RCFAD2	101	u	AFS	DYSIIDDVVC	LANDOCIMON	VECUUNCUE	150
NCT AUZ	101	M A W	AHUCGHHAF5	DYQLLDDVVG	LILHSCLLVP	YFSWKHSHRR	150
LFFAH12		100	170	תמו	100		150
FAH12	121	HHSNNGSLEK	DEVFVPPKKA	AVKWYVKYL-	NNPLGRILVL	THACTLAND	200
ATFAD2	7 7 1	いいつはまびつじたん	DEVENDEN	KICMACAAC"	MMDDCDVITI	A A T 1 A 1 A 1 A 1 A 1	200
BNFAD2	4 3 1	いいろはしなつしただ	DEVENDEN	AIKMYCVVI.	MMOLCOTMA	THACHARA	200
GMFAD2	131	UU SHI G SEEK	UEVEVER RS	DISSGTAST-	STTECOTUME	TUDETLOUGH	200
	- 171	いいつけいはつしいだ	DE A P A P K P K P	KANTECAA!	MMDICOAUCE	LVTITIGWPM	200
GMFAUZ-	- 171	UU 211 1 0 2 E E K	DEVENTEDES	C. LKMAZKAI -	MMDDCDVITA	AMTITIONS	
LMFAUZ	121	HHSNTGSLER	DEVFVPKKKE	AI PWYTPYVY	MNDVCDVVLI	VVOLTICHE	200
RCFAD2	151	HHSNTGSLER	DEVFVPKKKS	SIRWYSKYL-	NNPPGRIMTI	AVTLSLGWPL	200
. ==		210	220	220	240		200
LFFAH12	201	YLAFNYSGRP	YDG-FASHFF	PHARTEVEN	OLOTHICO.		
FAH12	-01	TEATHEOURP	IDK-FACHYII	PAUDIECEDE	DIDIVIANO	T C 1 T T C 11 1 1 1 1	250
ATFAD2		' - ~ ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '	IUUTTALMEE	PNAPIANNOE	DINIVICALA	* * * * * * * * * * * * * * * * * * * *	250
BNFAD2	-01	' LAINI JUKE	1 Duitzal Men	PNADIANUDE	DIATVICALA		250
GMFAD2 - 1		1 LACIN V 311 X P	TIIV-FAVHVE	DAYDIACHDE	DI 1 1 11 11 0 0 11 1		250
GMFAD2-2	201	YLALNYSGRP	YDR-FACHYD	PYCPIYCDE	WELLIA20AV	LESVIYSLYR	250
ZMFAD2	E Q 1	ILMINADURY	TPR-FALHEII	PACOLANUDE	DANTENCOAC	1111 4 44 4 6 6 4 4 4 4	250
RCFAD2	201	YLAFNVSGRP	YDR - FACHYD	DACDIANOUS	MAULEVSUAG	VVAVAFGLYK	250
		2	. SK TACHID	FIGELTNUKE	RIEIFISDAG	VLAVTFGLYQ	250

Figure 9A

, ,			c out	3 100	CCI	116	IA	1 (1)	4 GCC	: 111	F AA	T GTA	t TC	A GG	y Arg T AGA	202 1007
		· GA		110	. 661	ICA	LAI	11(. 110	: CC1	CAT	T GCA	CC1	T AT	e Phe C TTT	218 1055
		, cu,	י ערר	· cuc		LAG	AIA	N IAC	. AIC	TCA	GAT	GCT	GG1	ATI	e Leu ΓCTA	234 1103
	• • •		101	GGI	CII	IAC	CGI	IAC	GCI	GCT	TCA	CAA	GGA	TTO	Thr ACT	250 1151
		, ,,,,		GIC	IAI	GGA	ЬΙА	CCG	CII	ITG	ATA	GTG	AAC	TTT	Phe TTC	266 1199
•	3.0	.,,	, GIA	ACI	110	116	CAG	CAC	ACT	CAT	CCT	TCG	TTA	CCT	His	282 1247
				UAU	100	GAA	166	AII	AGA	GGA	GCT	Leu TTG	GTT	ACG	GTA	298 1295
		unc	171	GGA	AIA	116	AAC	AAG	GIG	TTC	CAT	Asn AAC	ATA	ACA	GAC	314 1343
		3.0	GCI	CAI	CAI	CIC	111	GCA	ACT	ATA	CCG	His CAT	TAT	AAC	GCA	330 1391
		ш.	ACA	UAU	ucu	AIA	AAG.	CCA	AIA	CII	GGT	Asp GAT	TAC	TAC	CAC	346 1439
		44.	ACA	cca	100	IAI	616	bcc	AIG	IAT	AGG	Glu GAA.	GCA	AAG	GAG	362 1487
Cys TGT	Leu CTC	Tyr TAT	Val GTA	Glu GAA	Pro CCG	Asp GAT	Thr ACG	Glu GAA	Arg CGT	Gly GGG	Lys AAG	Lys AAA	Gly GGT	Val GTC	Tyr TAC	378 1535
Tyr TAT	Tyr	Asn AAC	Asn AAT	Lys aag	Leu TTA	TGA	GGC	TGA	TAG	GGC	GAG	·AGA	AGT	GCA	ATT	384 1583
ATC	AAT	CTT	CAT	TTC	CAT	GTT	TTA	GGT	GTC	TTG	TTT	AAG	AAG	СТА	TĢC	1631
												TCT				1679
												CTG				1727
												ATA				1775
AAT	TGA	CCA	CAA	NAT	ATC	CAA .	AAC	CGG	CTA	TCC		TTC				1823
AAA	CCG	GAT	ATC	CAA	ATT	TCC	AGA	GTA	CTT	AG						1855

Figure 8B

AT	GAA	GCT	TTA	TAA	GAA	STT	AGT	777	CTC	TGG	TGA	CAG	AGA	AAT	TNT	47
GTC	AAT	TGG	TAG	TGA	CAG	TTG	AAG	CAA	CAG	GAA	CAA	CAA	GGA	TGG	TTG	95
GTG	NTG	ATG	CTG	÷TG	TGG	-GA	TGT	STT	ATT	CAT	CA÷	ATA	CTA	AAT	ACT	143
ACA	TTA	CTT	GTT	GCT	GCC	TAC	TTC	TCC	TAT	TTC	CTC	CGC	CAC	CCA	TTT	191
TGG	ACC	CAC	GAN	CCT	TCC	∴TT	TAA	ACC	СТС	TCT	CGT	GCT	ATT	CAC	CAG	239
AAG	AGA	AGC	CAA	GAG	AGA	GAG	AGA	GAG	AAT	GTT	CTG	AGG	ATC	ATT	GTC	287
TTC	TTC	ATC	GTT	ATT	AAC	GTA	AGT	TTT	TTT	TGA	CCY	СТС	ATA	TCT	AAA	335
ATC	TAG	TAC	ATG	CAA	TAG	ΑTT	AAT	GAC	TGT	TCC	TTC	TTT	TGA	TAT	TTT	383
CAG	CTT	стт	GAA	ттс	AAG	Met ATG	Gly GGT	Ala GCT	Gly	Gly GGA	Arg AGA	Ile	Met ATG	Val GTT	Thr ACC	10 431
Pro CCC	Ser TCT	Ser TCC	Lys AAG	Lys AAA	Ser	Glu GAA	Thr	Glu GAA	Ala GCC	Leu CTA	Lys AA;	Arg CGT	Gly GGA	Pro CCA	Cys TGT	26 479
Glu	Lys	Pro	Pro	Phe	Thr	Val	Lys	ASP	Leu	Lys	Lys	Ala	Ile	Pro	Gln	42
GAG	AAA	CCA	CCA	TTC	ACT	GTT	AAA	GAT	CTG	AAG	AAA		ATC	CCA	CAG	527
His	Cys	Phe	Lys	Arg	Ser	Ile	Pro	Arg	Ser	Phe	Ser	Tyr	Leu	Leu	Thr	58
CAT	TGT	TTC	AAG	CGC	TCT	ATC	CCT	CGT	TCT	TTC	TCC	TAC	CTT	CTC	ACA	575
Asp	Ile	Thr	Leu	Val	Ser	Cys	Phe	Tyr	Tyr	Val	Ala	Thr	Asn	Tyr	Phe	74
GAT	ATC	ACT	TTA	GTT	TCT	TGC	TTC	TAC		GTT	GCC	ACA	AAT	TAC	TTC	623
Ser	Leu	Leu	Pro	Gln	Pro	Leu	Ser	Thr	Tyr	Leu	Ala	Trp	Pro	Leu	Tyr	90
TCT	CTT	CTT	CCT	CAG	CCT	CTC	TCT	ACT	TAC	CTA	GCT	TGG	CCT	CTC	TAT	671
Trp	Val	Cys	Gln	Gly	Cys	Val	Leu	Thr	Gly	Ile	Trp	Val	Ile	Gly	His	106
TGG	GTA	TGT	CAA	GGC	TGT	GTC	TTA	ACC	GGT	ATC	TGG	GTC	ATT	GGC	CAT	719
Glu	Cys	Gly	His	His	Ala	Phe	Ser	Asp	Tyr	Gln	Trp	Val	Asp	Asp	Thr	122
GAA	TGT	GGT	CAC	CAT	GCA	TTC	AGT	GAC	Tat	CAA	TGG	GTA	GAT	GAC		767
Val	Gly	Phe	Ile	Phe	His	Ser	Phe	Leu	Leu	Val	Pro	Tyr	Phe	Ser	Trp	138
GTT	GGT	TTT	ATC	TTC	CAT	TCC	TTC	CTT	CTC	GTC	CCT	TAC	TTC	TCC	TGG	815
Lys	Tyr	Ser	His	Arg	Arg	His	His	Ser	Asn	Asn	Gly	Ser	Leu	Glu	Lys	154
AAA		AGT	CAT	CGT	CGT	CAC	CAT	TCC	AAC	AAT	GGA	TCT	CTC	GAG	AAA	863
Asp	Glu	Val	Phe	Val	Pro	Pro	Lys	Lys	Ala	Ala	V a 1	Lys	Trp	Tyr	Val	170
GAT	GAA	GTC	TTT	GTC	CCA	CCG	AAG	AAA	GCT	GCA	GTC	AAA	TGG	TAT	GTT	911
Lys	Tyr	Leu	Asn	Asn	Pro	Leu	Gly	Arg	Ile	Leu	V a 1	Leu	Thr	Val	Gìn	186
AAA	TAC	CTC	AAC	AAC	CCT	CTT	GGA	CGC	ATT	CTG	GTG	TTA	ACA	GTT	CAG	959

Figure 8A
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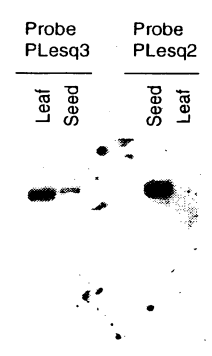


FIG.7

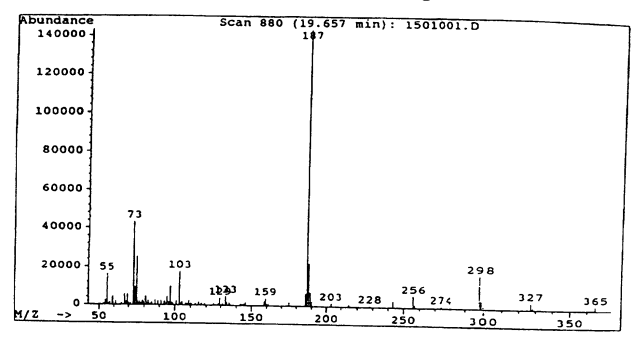
10	20	30	40	50	60
TATAGGCACC	GGAGGCACCA	TTCCAACACA	GGATCCCTCG	AAAGAGATGA	AGTATTTGTC
70	80	90	100	110	120
CCAAAGCAGA	AATCCGCAAT	CAAGTGGTAC	GGEGAATACC	TCAACAACCC	TCCTGGTCGC
130	140	150	160	170	180
ATCATGATGT	TAACTGTCCA	GTTCGTCCTC	GGATGGCCCT	TGTACTTAGC	CTTCAACGTT
190	200	210	220	230	240
TCTGGCAGAC	CCTACAATGG	TTTCGCTTCC	CATTTCTTCC	CCAATGCTCC	TATCTACAAC
250	260	270	280	290	300
GACCGTGAAC	GCCTCCAGAT	TTACATCTCT	GATGCTGGTA	TTCTAGCCGT	CTGTTATGGT
310	320	330	340	350	360
CTTTACCGTT	ACGCTGTTGC	ACAAGGACTA	GCCTCAATGA	TCTGTCTAAA	CGGAGTTCCG
370	380	390	400	410	420
CTTCTGATAG	TTAACTTTTT	CCTCGTCTTG	ATCACTTACT	TACAACACAC	TCACCCTGCG
430	440	450	460	470	480
TTGCCTCACT	ATGATTCATC	AGAGTGGGAT	TGGCTTAGAG	GAGCTTTAGC	TACTGTAGAC
490	500	510	520	530	540
AGAGACTATG	GAATCTTGAA	CAAGGTGTTC	CATAACATCA	CAGACACCCA	CGTCGCACAC
550					-
CACT					

Figure 6

10	20	30	40	50	60
TATTESCACC	GGCGGCACCA	TTCCAACAAT	GGATCCCTAG	AAAAAGATGA	AGTCTTTGTC
70	80	90	100	110	120
CCACCTAAGA	AAGCTGCAGT	CANATGGTAT	GTCAAATACC	TCAACAACCC	TCTTGGACGC
130	140	150	160	170	180
ATTCTGGTGT	TAACAGTTCA	GTTTATCCTC	GGGTGGCCTT	TGTATCTAGC	CTTTAATGTA
190	200	210	220	230	240
TCAGGTAGAC	CTTATGATGG	TTTCGCTTCA	CATTTCTTCC	CTCATGCACC	TATCTTTAAG
250	260	270	280	290	300
GACCGTGAAC	GTCTCCAGAT	ATACATCTCA	GATGCTGGTA	TTCTAGCTGT	CTGTTATGGT
310	320	330	340	350	360
CTTT+CCGTT	ACGCTGCTTC	ACAAGGATTG	ACTGCTATGA	TCTGCGTCTA	CGGAGTACCG
370	380	390	400	410	420
CTTTTGATAG	TGAACTTTTT	CCTTGTCTTG	GTCACTTICT	TGCAGCACAC	TCATCCTTCA
430	440	4 50	460	470	480
TTACCTCACT	ATGATTCAAC	CGAGTGGGAA	TGGATTAGAG	GAGCTTTGGT	TACGGTAGAC
490	500	510	520	530	540
AGAGACTATG	GAATCTTGAA	CAAGGTGTTT	CACAACATAA	CAGACACCCA	CGTAGCACAC
550					
CAC					

Figure 5

4C Mass spectrum of peak 12 from figure 3B



4D Mass spectrum of peak 13 from figure 3B

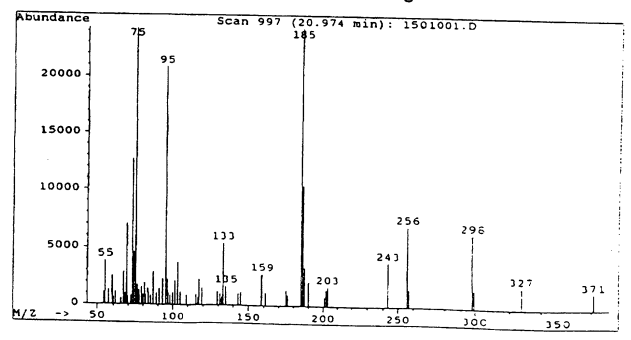
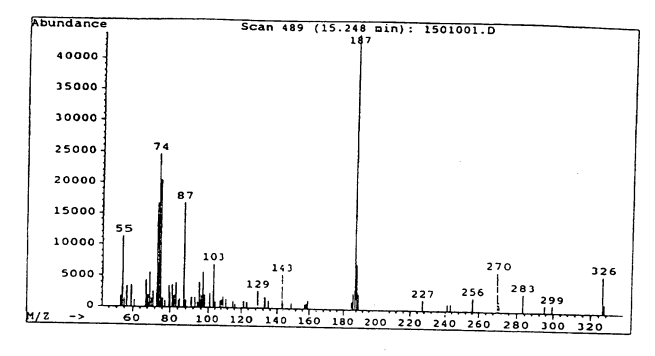


Figure 4A Mass spectrum of peak 10 from figure 3B



4B Mass spectrum of peak 11 from figure 3B

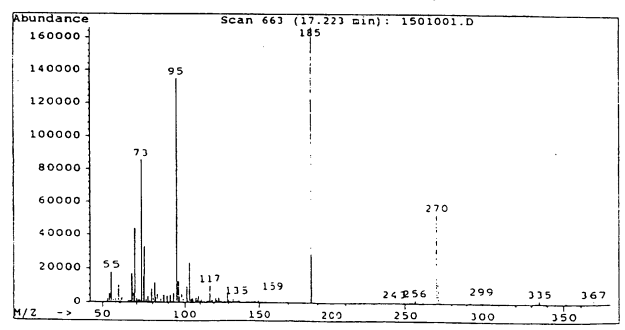
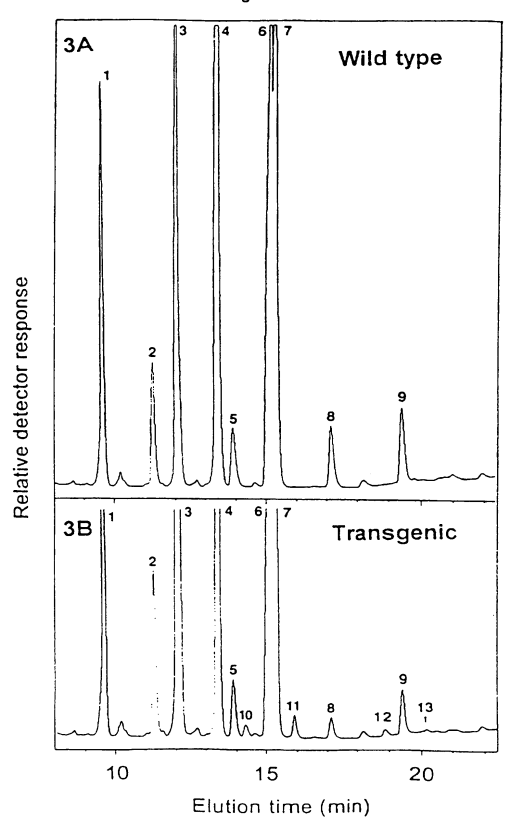


Figure 3



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lon #2: Mass 299

ion #3: Mass 270 (characteristic rearrangement ion)

Ion #4: Mass 185 (desaturated analog of Ion #1)

ion #5: Mass 298 (elongated analog of ion #3)

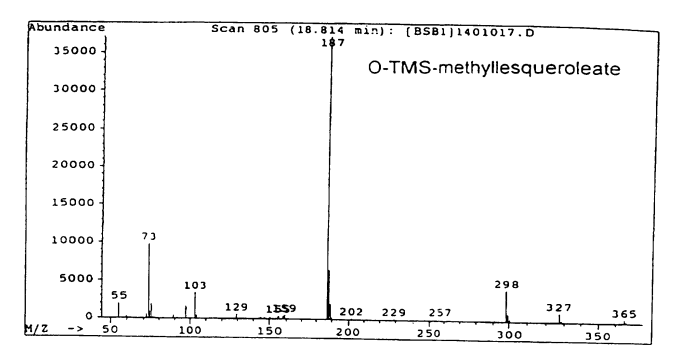
Ion #6: Mass 327 (elongated analog of ion

$$(CH_3)_3$$
-Si-O-CH-CH₂-CH=CH-(CH₂)₉-C-O-CH₃ +

Figure 2

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1C



1D

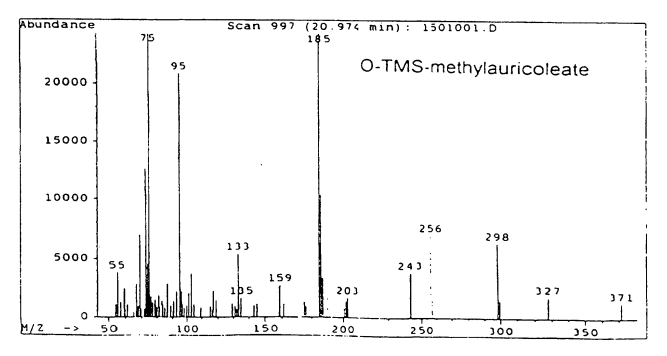
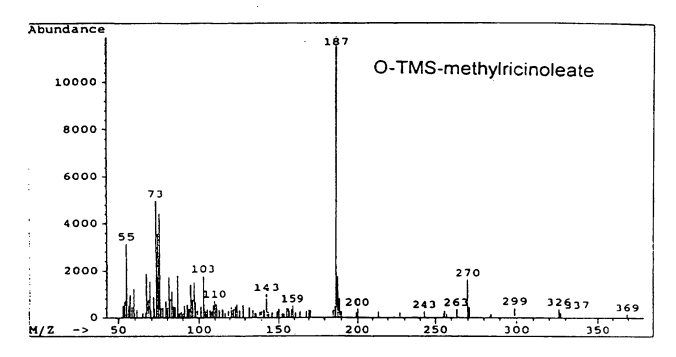
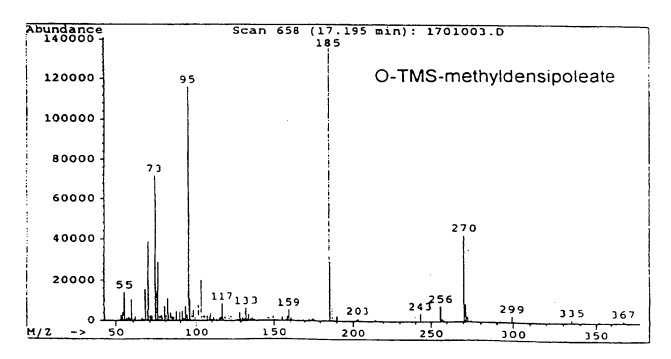


Figure 1A



1B



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- 25. The recombinant nucleic acid of Claim 23, wherein said sequence is obtainable from Lesquerella fendleri.
- 26. The recombinant nucleic acid of Claim 23, wherein said nucleic acid contains a sequence encoding at least one of fatty acid desaturase or fatty acid hydroxylase in which one or more essential histidine residues have been mutated.
- 27. The method of Claim 1 further comprising: processing the seed containing the altered amount of the unsaturated fatty acid to obtain oil and/or seed meal.
 - 28. Oil obtained by the method of Claim 27.
- 29. Seed meal obtained by the method of Claim 27.
 - 30. Plant obtained by the method of Claim 1.
- 31. The method of Claim 11 further comprising: processing the seed containing the altered amount of the unsaturated fatty acid to obtain oil and/or seed meal.
 - 32. Oil obtained by the method of Claim 31.
- 33. Seed meal obtained by the method of Claim 31.
 - 34. Plant obtained by the method of Claim 11.

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amino acid identity of 60% or greater to SEQ ID NO:4.

- 18. The recombinant nucleic acid of Claim 17, wherein the amino acid identity is 90% or greater to SEQ ID NO:4.
- 19. The recombinant nucleic acid of Claim 17, wherein the amino acid identity is 100% of SEQ ID NO:4.
- 20. The recombinant nucleic acid of Claim 17, wherein said nucleic acid contains a sequence having a nucleotide identity of 90% or greater to SEQ ID NO:1, 2 or 3.
- 21. The recombinant nucleic acid of Claim 17, wherein said nucleic acid contains SEQ ID NO:1, 2 or 3.
- 22. The recombinant nucleic acid of Claim 17, wherein said sequence is obtainable from a plant species producing a hydroxylated fatty acid.
- 23. A recombinant nucleic acid suitable for use in Claim 1, wherein said nucleic acid contains a sequence encoding at least one of fatty acid desaturase or fatty acid hydroxylase.
- 24. The recombinant nucleic acid of Claim 23, wherein said sequence is obtainable from Ricinus communis (L.) (castor).

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- 12. The method of Claim 11, wherein said nucleic acid contains a sequence which encodes the dominant negative mutant of fatty acid hydroxylase in which one or more essential histidine residues have been mutated.
- 13. The method of Claim 11, wherein said nucleic acid contains a sequence which encodes the dominant negative mutant of fatty acid hydroxylase which thereby alters the amount of the unsaturated fatty acid in the seed.
- 14. The method of Claim 11, wherein said nucleic acid contains a sequence which encodes the dominant negative mutant of fatty acid desaturase in which one or more essential histidine residues have been mutated.
- 15. The method of Claim 11, wherein said nucleic acid contains a sequence which encodes the dominant negative mutant of fatty acid desaturase which thereby alters the amount of the unsaturated fatty acid in the seed.
- 16. The method of Claim 11, wherein said plant is selected from the group consisting of rapeseed, Crambe, Brassica juncea, canola, flax, sunflower, safflower, cotton, cuphea, soybean, peanut, coconut, oil palm and corn.
- 17. A recombinant nucleic acid suitable for use in Claim 1, wherein said nucleic acid contains a sequence encoding a fatty acid hydroxylase with an

- 7. The method of Claim 1, wherein said plant is transformed with a nucleic acid containing a sequence which encodes a fatty acid desaturase or derivative thereof.
- 8. The method of Claim 7, wherein said derivative is a dominant negative mutant which thereby alters the amount of the unsaturated fatty acid in the seed.
- 9. The method of Claim 7, wherein said derivative is a mutant fatty acid desaturase in which one or more essential histidine residues have been mutated which thereby alters the amount of the unsaturated fatty acid in the seed.
- 10. The method of Claim 1, wherein said plant is selected from the group consisting of rapeseed, Crambe, Brassica juncea, canola, flax, sunflower, safflower, cotton, cuphea, soybean, peanut, coconut, oil palm and corn.
- 11. A method of altering an amount of a unsaturated fatty acid comprising:
- (a) transforming a plant cell with a nucleic acid containing a sequence which encodes a fatty acid hydroxylase or a dominant negative mutant of fatty acid hydroxylase or a dominant negative mutant of fatty acid desaturase,
- (b) growing a seed-bearing plant from the transformed plant cell of step (a), and
- (c) identifying a seed from the plant of step (b) with the altered amount of the unsaturated fatty acid in the seed.

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WHAT IS CLAIMED IS:

- 1. A method of altering an amount of an unsaturated fatty acid in a seed of a plant comprising: decreasing a fatty acid desaturase activity in the seed by genetic manipulation of at least one of fatty acid desaturase or fatty acid hydroxylase.
- 2. The method of Claim 1, wherein an endogenous gene for said fatty acid hydroxylase is mutated and thereby decreases fatty acid hydroxylase activity in the seed.
- 3. The method of Claim 1, wherein said plant is transformed with a nucleic acid containing a sequence which encodes a fatty acid hydroxylase or derivative thereof.
- 4. The method of Claim 3, wherein said derivative is a dominant negative mutant which thereby alters the amount of the unsaturated fatty acid in the seed.
- 5. The method of Claim 3, wherein said derivative is a mutant fatty acid hydroxylase in which one or more essential histidine residues have been mutated which thereby alters the amount of the unsaturated fatty acid in the seed.
- 6. The method of Claim 1, wherein an endogenous gene for said fatty acid desaturase is mutated and thereby decreases fatty acid desaturase activity in the seed.

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Phe Thr Met Pro	Phe	Thr	Met	Dro
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(2)	INFORMATION FOR SEQ ID NO:12	
	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 nucleotides (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:12:	
GCTCT	TTTTGT GCGCTCATTC	20
(2)	INFORMATION FOR SEQ ID NO:13	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 nucleotides (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:13:	
CGGT	ACCAGA AAACGCCTTG	20
(2)	INFORMATION FOR SEQ ID NO:14	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 nucleotides (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:14:	
TAYWS	SNCAYM GNMGNCAYCA	20
(2)	INFORMATION FOR SEQ ID NO:15	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 nucleotides (B) TYPE: nucleotide (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:15:	
RTGR'	TGNGCN ACRTGNGTRT C	21

Leu	Glu	Arg	Asp	Glu 55	Val	Phe	Val	Pro	Lys 60
Lys	Lys	Ser	Ser	Ile 65	Arg	Trp	Tyr	Ser	Lys 70
Tyr	Leu	Asn	Asn	Pro 75	Pro	Gly	Arg	Ile	Met 80
Thr	Ile	Ala	Val	Thr 85	Leu	Ser	Leu	Gly	Trp 90
Pro	Leu	Tyr	Leu	Ala 95	Phe	Asn	Val	Ser	Gly 100
Arg	Pro	Tyr	Asp	Arg 105	Phe	Ala	Cys	His	Tyr 110
Asp	Pro	Tyr	Gly	Pro 115	Ile	Tyr	Asn	Asp	Arg 120
Glu	Arg	Ile	Glu	Ile 125	Phe	Ile	Ser	Asp	Ala 130
Gly	Val	Leu	Ala	Val 135	Thr	Phe	Gly	Leu	Tyr 140
Gln	Leu	Ala	Ile	Ala 145	Lys	Gly	Leu	Ala	Trp 150
Val	Val	Cys	Val	Tyr 155	Gly	Val	Pro	Leu	Leu 160
Val	Val	Asn	Ser	Phe 165	Leu	Val	Leu	Ile	Thr 170
Phe	Leu	Gln	His	Thr 175	His	Pro	Ala	Leu	Pro 180
His	Tyr	Asp	Ser	Ser 185	Glu	Trp	Asp	Trp	Leu 190
Arg	Gly	Ala	Leu	Ala 195	Thr	Val	Asp	Arg	Asp 200
Tyr	Gly	Ile	Leu	Asn 205	Lys	Val	Phe	His	Asn 210
Ile	Thr	Asp	Thr	Gln 215	Val	Ala	His	His	Leu 220

Asp Tyr Gly Ile Leu Asn Arg Val Phe His 300
Asn Ile Thr Asp Thr His Val Ala His His 310
Leu Phe Ser Thr Met Pro His Tyr His Ala 320
Met Glu Ala Thr Lys Ala Ile Arg Pro Ile 330
Leu Gly Asp Tyr Tyr His Phe Asp Pro Thr 340
Pro Val Ala Lys Ala Thr Trp Arg Glu Ala 350
Gly Glu Cys Ile Tyr Val Glu Pro Glu Asp 360
Arg Lys Gly Val Phe Trp Tyr Asn Lys Lys 370
Phe Xaa

- (2) INFORMATION FOR SEQ ID NO:11
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 224 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Trp Val Met Ala His Asp Cys Gly His His
5

Ala Phe Ser Asp Tyr Gln Leu Leu Asp Asp

Val Val Gly Leu Ile Leu His Ser Cys Leu

Leu Val Pro Tyr Phe Ser Trp Lys His Ser 35

His Arg Arg His His Ser Asn Thr Gly Ser

Met	Val	Pro	Tyr	Phe 125	Ser	Trp	Lys	Tyr	Ser 130
His	Arg	Arg	His	His 135	Ser	Asn	Thr	Gly	Ser 140
Leu	Glu	Arg	Asp	Glu 145	Val	Phe	Val	Pro	Lys 150
Lys	Lys	Glu	Ala	Leu 155	Pro	Trp	Tyr	Thr	Pro 160
Tyr	Val	Tyr	Asn	Asn 165	Pro	Val	Gly	Arg	Val 170
Val	His	Ile	Val	Val 175	Gln	Leu	Thr	Leu	Gly 180
Trp	Pro	Leu	Tyr	Leu 185	Ala	Thr	Asn	Ala	Ser 190
Gly	Arg	Pro	Tyr	Pro 195	Arg	Phe	Ala	Cys	His 200
Phe	Asp	Pro	Tyr	Gly 205	Pro	Ile	Tyr	Asn	Asp 210
Arg	Glu	Arg	Ala	Gln 215	Ile	Phe	Val	Ser	Asp 220
Ala	Gly	Val	Val	Ala 225	Val	Ala	Phe	Gly	Leu 230
Tyr	Lys	Leu	Ala	Ala 235	Ala	Phe	Gly	Val	Trp 240
Trp	Val	Val	Arg	Val 245	Tyr	Ala	Val	Pro	Leu 250
Leu	Ile	Val	Asn	Ala 255	Trp	Leu	Val	Leu	Ile 260
Thr	Tyr	Leu	Gln	His 265	Thr	His	Pro	Ser	Leu 270
Pro	His	Tyr	Asp	Ser 275	Ser	Glu	Trp	Asp	Trp 280
Leu	Arg	Gly	Ala	Leu 285	Ala	Thr	Met	Asp	Arg 290

Trp Tyr Asn Asn Lys Leu Ala Met Glu Ala 295 300

Thr Val

- (2) INFORMATION FOR SEQ ID NO:10
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 372 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Met Gly Ala Gly Gly Arg Met Thr Glu Lys 5
- Glu Arg Glu Lys Gln Glu Gln Leu Ala Arg
 15 20
- Ala Thr Gly Gly Ala Ala Met Gln Arg Ser 25 30
- Pro Val Glu Lys Pro Pro Phe Thr Leu Gly
- Gln Ile Lys Lys Ala Ile Pro Pro His Cys
- Phe Glu Arg Ser Val Leu Lys Ser Phe Ser 55 60
- Tyr Val Val His Asp Leu Val Ile Ala Ala 65 70
- Ala Leu Leu Tyr Phe Ala Leu Ala Ile Ile
 75 80
- Pro Ala Leu Pro Ser Pro Leu Arg Tyr Ala 85 90
- Ala Trp Pro Leu Tyr Trp Ile Ala Gln Gly 95 100
- Ala Phe Ser Asp Tyr Ser Leu Leu Asp Asp 105
- Val Val Gly Leu Val Leu His Ser Ser Leu 115 120

Thr	Leu	Gly	Trp	Pro 125	Leu	Tyr	Leu	Ala	Leu 130
Asn	Val	Ser	Gly	Arg 135	Pro	Tyr	Asp	Arg	Phe 140
Ala	Cys	His	Tyr	Asp 145	Pro	Tyr	Gly	Pro	Ile 150
Tyr	Ser	Val	Ile	Ser 155	Asp	Ala	Gly	Val	Leu 160
Ala	Val	Val	Tyr	Gly 165	Leu	Phe	Arg	Leu	Ala 170
Met	Ala	Lys	Gly	Leu 175	Ala	Trp	Val	Val	Cys 180
Val	Tyr	Gly	Val	Pro 185	Leu	Leu	Val	Val	Asn 190
Gly	Phe	Leu	Val	Leu 195	Ile	Thr	Phe	Leu	Gln 200
His	Thr	His	Val	Ser 205	Glu	Trp	Asp	Trp	Leu 210
Arg	Gly	Ala	Leu	Ala 215	Thr	Val	Asp	Arg	Asp 220
Tyr	Gly	Ile	Leu	Asn 225	Lys	Val	Phe	His	Asn 230
Ile	Thr	. Asp	Thr	His 235	Val	Ala	His	His	Leu 240
Phe	e Sei	Thr	Met	Pro 245	His	Tyr	His	Ala	Met 250
Glu	ı Ala	a Thr	. Val	Glu 255	Tyr	Tyr	Arg	Phe	Asp 260
Gl	ı Th	r Pro	o Phe	e Val	Lys	s Ala	Met	Trp	Arg 270
Gl	u Al	a Ar	g Gli	ц Суя 27!	s Ile	е Туг	val	l Glu	280
As	p Gl	n Se	r Th	r Gl 28	u Se	r Lys	s Gly	y Val	290

Asp Glu Gly Thr Ser Glu Lys Gly Val Tyr 295 300

Trp Tyr Arg Asn Lys Tyr Leu Arg Val

- (2) INFORMATION FOR SEQ ID NO:9
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 302 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Phe Ser Tyr Val Val Tyr Asp Leu Thr Ile
5

Ala Phe Cys Leu Tyr Tyr Val Ala Thr His
15 20

Tyr Phe His Leu Leu Pro Gly Pro Leu Ser 25 30

Phe Arg Gly Met Ala Ile Tyr Trp Ala Val

Gln Gly Cys Ile Leu Thr Gly Val Trp Val
45
50

Val Ala Phe Ser Asp Tyr Gln Leu Leu Asp 55 60

Asp Ile Val Gly Leu Ile Leu His Ser Ala 65 70

Leu Leu Val Pro Tyr Phe Ser Trp Lys Tyr
75 80

Ser His Arg Arg His His Ser Asn Thr Gly 85 90

Ser Leu Glu Arg Asp Glu Val Phe Val Pro 95

Lys Val Ser Lys Tyr Leu Asn Asn Pro Pro 105 110

Gly Arg Val Leu Thr Leu Ala Val Thr Leu 115 120

Ser	Leu	Leu	Val	Thr 125	Leu	Thr	Ile	Gly	Trp 130
Pro	Met	Tyr	Leu	Ala 135	Phe	Asn	Val	Ser	Gly 140
Arg	Pro	Tyr	Asp	Ser 145	Phe	Ala	Ser	His	Tyr 150
His	Pro	Tyr	Arg	Val 155	Arg	Leu	Leu	Ile	Tyr 160
Val	Ser	Asp	Val	Ala 165	Leu	Phe	Ser	Val	Thr 170
Tyr	Ser	Leu	Tyr	Arg 175	Val	Ala	Thr	Leu	Lys 180
Gly	Leu	Val	Trp	Leu 185	Leu	Cys	Val	Tyr	Gly 190
Val	Pro	Leu	Leu	Ile 195	Val	Asn	Gly	Phe	Leu 200
Val	Thr	Ile	Thr	Tyr 205	Leu	Arg	Val	His	Tyr 210
Asp	Ser	Ser	Glu	Trp 215	Asp	Trp	Leu	Lys	Gly 220
Ala	Leu	Ala	Thr	Met 225	Asp	Arg	Asp	Tyr	Gly 230
Ile	Leu	Asn	Lys	Val 235	Phe	His	His	Ile	Thr 240
Asp	Thr	His	Val	Ala 245	His	His	Leu	Phe	Ser 250
Thr	Met	Pro	His	Tyr 255	His	Leu	Arg	Val	Lys 260
Pro	Ile	Leu	Gly	Glu 265	-Tyr	Tyr	Gln	Phe	Asp 270
Asp	Thr	Pro	Phe	Tyr 275	Lys	Ala	Leu	Trp	Arg 280
Glu	Ala	Arg	Glu	Cys 285	Leu	Tyr	Val	Glu	Pro 290

Gly Glu Lys Lys Gly Val Phe Trp Tyr Asn 375

Asn Lys Leu Xaa

- (2) INFORMATION FOR SEQ ID NO:8
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 309 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
- Ser Leu Leu Thr Ser Phe Ser Tyr Val Val
 5
- Tyr Asp Leu Ser Phe Ala Phe Ile Phe Tyr
 15 20
- Ile Ala Thr Thr Tyr Phe His Leu Leu Pro 25 30
- Gln Pro Phe Ser Leu Ile Ala Trp Pro Ile 35 40
- Tyr Trp Val Leu Gln Gly Cys Leu Leu Thr
 45 50
- Arg Val Cys Gly His His Ala Phe Ser Lys
- Tyr Gln Trp Val Asp Asp Val Val Gly Leu
 65 70
- Thr Leu His Ser Thr Leu Leu Val Pro Tyr
 75 80
- Phe Ser Trp Lys Ile Ser His Arg Arg His 85 90
- His Ser Asn Thr Gly Ser Leu Asp Arg Asp 95
- Glu Arg Val Lys Val Ala Trp Phe Ser Lys
 105 110
- Tyr Leu Asn Asn Pro Leu Gly Arg Ala Val